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(54) Title: IMPROVED ADENOVIRAL VECTORS AND PRODUCER CELLS

(57) Abstract

An adenoviral vector wherein the adenoviral genome has been modified to reduce the host immune and inflammatory responses to the vector. In one embodiment, the adenoviral vector including an adenoviral 5' ITR; an adenoviral 3' ITR; an adenoviral encapsidation signal; at least one DNA sequence encoding a protein or polypeptide of interest; and a promoter controlling the DNA sequence(s) encoding the protein(s) or polypeptide(s) of interest. The adenoviral vector is free of all or a portion of each of the adenoviral E1 and E4 DNA sequences, or is free of all or a portion of each of the adenoviral E1 and E2 DNA sequences, or is free of all or a portion of each of the E1. E2 and E4 DNA sequences. Also disclosed and claimed is a producer cell for generating adenoviral vector particles, wherein the producer cell includes an adenoviral E1 DNA sequence and an adenoviral E4 DNA sequence, or includes an adenoviral E1 DNA sequence and an adenoviral E2a DNA sequence, or includes the adenoviral E1, E2a and E4 DNA sequences.

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IMPROVED ADENOVIRAL VECTORS AND PRODUCER CELLS

This application is a continuation-in-part of application

Serial No. 08/355,087, filed December 12, 1994.

This invention relates to adenoviral vectors wherein the adenoviral genome has been modified to reduce the host immune and inflammatory responses to the vector. More particularly, this invention relates to adenoviral vectors which are free of all or a part of the adenoviral E1 and E4 DNA sequences, or are free of all or a part of the adenoviral E1 and E2 DNA sequences or are free of all or a portion of each of the E1, E2, and E4 regions and further may or may not contain some of the E3 region genes operationally linked to a functional promoter, and to producer cells for generating adenoviral particles from such vectors.

BACKGROUND OF THE INVENTION

Adenovirus genomes are linear, double-stranded DNA molecules about 36 kilobase pairs long. Each extremity of the viral genome has a short sequence known as the inverted terminal repeat (or ITR), which is necessary for viral The well-characterized molecular genetics of replication. an advantageous vector for gene adenovirus render it The knowledge of the genetic organization of transfer. adenoviruses allows substitution of large fragments of viral In addition, recombinant DNA with foreign sequences. adenoviruses are structurally stable and no rearranged viruses are observed after extensive amplification.

WO 96/18418 PCT/US95/15947.

Adenoviruses thus may be employed as delivery vehicles for introducing desired genes into eukaryotic cells, whereby the adenovirus delivers such genes to eukaryotic cells by binding cellular receptors, internalizing via coated pits, disrupting endosomes, and releasing particles to the cytoplasm followed by nuclear translocation and molecular expression of the adenovirus genetic program.

In general, an adenovirus genome includes the El region, the B2 region, the E3 region, the B4 region, and the major late region as well as several other minor regions. region provides functions associated with transformation, transcription upregulation, and control of replication. E2 region is essential for DNA replication, and other viral functions including transcriptional regulation. region provides for the adenovirus' defense against the host's immune system. The E3 region encodes proteins which inhibit the host cell's ability to present viral proteins as The E4 region is essential for several foreign antigens. important viral functions, including control of late gene expression, shutting down host cell function by the virus, The major late region encodes the and virus replication. major adenoviral structural proteins.

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Adenoviral vectors have been constructed in which at least a majority of the E1 and E3 sequences have been The B1 deletion renders the vector replication defective, the B3 region deletion provides space for insertion of the desired gene into the adenoviral vector without exceeding the maximum acceptable size Applicants have found that when such packageable genome. vectors have been placed into target cells or organs in vivo, however, a sharp decrease in expression of the desired gene results at about 1 to 3 weeks post-infection. In addition, it was found that an inflammatory response may develop in a target organ, and that the adenovirus vector may induce a cytotoxic T-lymphocyte (CTL) response directed against the

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vector-transcored host cells. This cresponse has the effect of eliminating the book vector transcored cells from the host may develop a newtralizing antibody response to such adenoviral vectors.

The is therefore an object of the present invention to provide an adenoviral vector which minimizes host response to the vector and increases the duration of vector persistence and expression, and to provide producer cell lines which will generate adenoviral particles from such vectors.

BRIEF DESCRIPTION OF THE DRAWINGS

The invention now will be described with respect to the drawings, wherein:

Figure 1 is a schematic of the construction of plasmid pSE280-E1;

Figure 2 is a map of plasmid pMAMneo-Luc;

Figure 3 is a schematic of the construction of plasmid pMAMneo-E1;

Figure 4 is a schematic of the construction of plasmid pGRE5-E1 from pSE280-E1 and pGRE5-2/EBV;

Figure 5 is a map of plasmid pGRB5-E1;

Figure 6 a schematic of the construction of plasmid pMAMneo-E2A;

Figure 7 is a map of plasmid pSE380;

Figure 8 is a schematic of the construction of plasmids pSE380-E4 and pSE380-ITR/E4;

Figure 9 is a schematic of the construction of plasmid pSE380GRE5/E4;

Figure 10 is a map of plasmid pTZ18R;

Figure 11 is a schematic of the construction of plasmids pTZE2A and pTZdlE2A;

Figure 12 is a schematic of the construction of plasmid pSE380-E3+;

Figure 13 is a schematic of the construction of plasmids pSE380-E3*E4* and pTZE3*E4*;

WO 96/15418 PCT/US95/15947*

Figure 14 is a schematic of the construction of plasmids pTZE3^{E4+}, pSE280E3^{E4+}, and pSPORT-1 E3^{E4+};

Figure 15 is a map of plasmid pSPORT-1;

Figure 16 is a schematic of the construction of plasmid pSPORT-1E2AE3E4+;

Figure 17 is a map of plasmid pBR322;

Figure 18 is a schematic of the construction of plasmids pBR322-E2AE3E4+ and pBRAd5-E2AE3E4+;

Figure 19 is a schematic of the construction of plasmids pSE380-E4 and pSE380E3*E4;

Figure 20 is a schematic of the construction of plasmid psE380E3+E4-(2);

Figure 21 is a schematic of the construction of plasmid pSE380E3E4'(2);

Figure 22 is a schematic of the construction of plasmid pspORT1/E2AE3E4 (2);

Figure 23 is a schematic of the production of an adenoviral vector having the genotype E1⁺, E2A⁻, E3⁻, E4⁺;

Figure 24 is a map of pAvS6;

Figure 25 is a schematic of the construction of an adenoviral vector having the genotype E1, E2A, E3, E4⁺ by homologous recombination;

Figure 26 is a schematic of the production of an adenoviral vector having the genotype E1, E2A, E3, E4⁺ by <u>in vitro</u> ligation and transfection;

Figure 27 is a schematic of the production of an adenoviral vector having the genotype E1, E2A, E3, E4 by homologous recombination;

Figure 28 is a schematic of the vector Av3nLacZ;

Figure 29 is a schematic of the production of an adenoviral vector having the genotype R1+, R4-;

Figure 30 is a schematic of the production of an adenoviral vector having the genotype E1, E2⁺, E3⁺, E4 by homologous recombination;

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Figure 31 is a schematic of the production of an adenoviral vector having the genotype E1, E2⁺, E3⁺, E4 by <u>in vitro</u> ligation and transfection;

Figure 32 is a schematic of the generation of Av4nLacZ by homologous recombination;

Figure 33 is a schematic of the generation of Av4nLacZ by in vitro ligation;

Figure 34 is a schematic of the generation of an Av4 (fourth generation adenovirus) type vector containing a transgene;

Figure 35 is a schematic of the construction of plasmid pAvS6-CFTR;

Pigure 36 is an immunoprecipitation blot for adenoviral hexon expression in A549 cells infected with an Av1 type vector (having a deletion of all or part of the E1 region), an Av2 vector (having a deletion of all or part of the E1 region and a temperature-sensitive mutation in the E2a gene), or an Av3 vector or Ad dl327;

Figure 37 is an immunoprecipitation blot for adenoviral hexon expression in E1+E2+ cells or E1+ cells infected with an Av1 vector, an Av2 vector, or an Av3 vector; and

Figure 38 is a blot showing adenovirus vector DNA replication in $E1^+E2^+$ or $E1^+$ cells infected with an Av1 vector, an Av2 vector, or an Av3 vector.

SUMMARY OF THE INVENTION

286

WO 96/18418 - 🐧 🔩

The present invention provides for an improved adenoviral vector wherein the adenoviral genome has been modified to reduce the host immune and inflammatory response to the vector. Such vector has a reduced viral DNA replication potential and reduced expression of adenoviral genes (e.g., hexon) in comparison to existing adenoviral vectors. Such modification(s) to the adenoviral genome may be effected through deletion(s) and/or mutation(s) of portion(s) of the adenoviral genome.

DETAILED DESCRIPTION OF THE INVENTION

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The adenoviral vectors described herein include several general types or generations, each of which are described These vectors include an adenovirus DNA genome containing an adenovirus 5'ITR; an adenovirus 3'ITR; adenovirus encapsidation signal; at least one DNA sequence encoding a protein or polypeptide of interest; and a promoter controlling the at least one DNA sequence encoding a protein or polypeptide of interest. One generation of adenoviral vectors (Av3 vectors) contain a deletion of all or part of the E1 region and also a deletion of all or a part of the E2 region (preferably all or part of the E2a region). A second generation of adenoviral vectors (Av4 vectors) contain a deletion of all or part of the E1 region and a deletion of all or a part of the E2 region (preferably all or part of the E2a region), and a deletion of all or a part of the E4 region. A third generation of adenoviral vectors (Av5 vectors) contain a deletion of all or a part of the E1 region, and a deletion of all or part of the E4 region. Each of these vectors may or may not contain some part of the E3 region operationally linked to a promoter which controls its expression.

In these vectors, the DNA sequence encoding a protein or polypeptide of interest, which is linked operationally to a promoter controlling its expression is sometimes hereinafter referred to as a transgene. Cell lines containing adenovirus structural genes under the control of an inducible promoter (i) are referred to as iE1 (E1 genes); iE2a (E2a genes); iE1/E2 (E1 and E2 genes); (i)E1/E4 (E1 and E4 genes); or iE1/E2a/E4 (E1, E2, and E4 genes).

In one embodiment, there is provided an adenoviral vector which includes an adenoviral 5' ITR; an adenoviral 3' ITR; an adenoviral encapsidation signal; at least one DNA sequence encoding a protein or polypeptide of interest; and a promoter controlling the DNA sequence(s) encoding a protein(s) or polypeptide(s) of interest. The adenoviral

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vector is free of all or a portion of the adenoviral E1 (preferably at least all or a portion of E1a) and E4 DNA sequences. In a preferred embodiment, the adenoviral vector also is free of all or a portion of adenoviral E1b DNA sequence. Such a vector is sometimes referred to as an adenoviral E1E4 vector.

The term "portion" as used herein means that a portion of the DNA sequence of a region (e.g., E1, E2, E3, and/or E4) of the adenoviral genome is removed such that the function of such region is eliminated or impaired. For example, a portion of the DNA sequence of a region of the adenoviral genome is removed such that one or more proteins normally encoded by such region are not expressed, or are expressed as a structure which eliminates or impairs the function of such proteins.

The determination of the amount of the DNA sequence of a region which is to be removed is determined by routine experimentation by one skilled in the art given the teachings herein. Preferably, all or substantially all of a coding region is removed.

In one preferred embodiment, at least open reading frame 6 (ORF 6) of the B4 DNA sequence is deleted. In another embodiment, at least open reading frames 3 and 6 of the B4 DNA sequence are deleted.

In another embodiment, there is provided an adenoviral vector which includes an adenoviral 5' ITR; an adenoviral 3' ITR; an adenoviral encapsidation signal; at least one DNA sequence encoding a protein or polypeptide of interest; and a promoter controlling the DNA sequence(s) encoding a protein(s) or polypeptide(s) of interest. The adenoviral vector is free of all or a portion of the adenoviral B1 and all or a portion of B2 (preferably at least all or a portion of B2a) DNA sequences. In a preferred embodiment, the adenoviral vector also is free of all or a portion of the

WO 96/18418 PCT/US95/15947

adenoviral E1b DNA sequence. Such vector is sometimes referred to as an adenoviral E1E2 vector.

In another embodiment, all or a portion of the E2b DNA sequence is deleted. In yet another embodiment, all or a portion of the E2a and E2b DNA sequences are deleted.

In another preferred embodiment, there is provided an adenoviral vector which includes an adenoviral 5' ITR; an adenoviral 3' ITR; an adenoviral encapsidation signal; at least one DNA sequence encoding a protein or polypeptide of interest; and a promoter controlling the DNA sequence(s) encoding the protein(s) or polypeptide(s) of interest. The adenoviral vector is free of all or a portion of the adenoviral E1 (including E1a and E1b), E2, and E4 DNA sequences. Such vector is sometimes referred to as an adenoviral E1E2E4 vector. Preferably, such vector is free of at least open reading frame 6 of the E4 DNA sequence.

In one preferred embodiment the vector is free of all or a portion of the B2a DNA sequence. Such vector is sometimes referred to as an adenoviral B1B2aB4 vector. In another embodiment, the vector is free of all or a portion of the B2b DNA sequence.

In one embodiment, the adenoviral vectors hereinabove described also include at least a portion of the adenoviral E3 DNA sequence, operationally linked to a promoter which allows expression of E3 region genes in the transduced host cells, and which enables the adenovirus to retain some or all of its immune defense system. The amount of the E3 DNA sequence included in the adenoviral vector is, in general, an amount of the E3 DNA sequence which will enable the adenoviral vector to evade or disable the host's immune response. Preferably, all of the genes of the E3 region are included in the vector, except the gene encoding the 11.6 Kda protein that causes cell lysis.

In another embodiment, the above measured adenoviral vectors do not contain the E3 region genes.

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The vectors of the present invention in a preferred embodiment are adenoviral particles wherein the genetic material contained in the virus lacks the genetic material which is specified as having been deleted.

Thus, in accordance with a preferred embodiment of the invention, the genome of the adenoviral vector contains the early regions (E1, E2, E3 and E4) and the major late regions to the extent necessary not to impair the function of the proteins encoded thereby, except for all or a portion of the regions which are specified as having been deleted from the genetic material of the adenovirus. Thus, if an adenoviral vector is defined as being free of all or a portion of E1 and E2, then such vector would contain the E3 and E4 regions to the extent necessary not to disrupt or substantially impair the function of the protein encoded thereby. All other regions, including the major late regions and minor transcriptional units, would also be present to the extent necessary not to disrupt or substantially impair the function of the proteins encoded thereby.

The adenoviral vector may be derived from any known adenovirus serotype, including, but not limited to, Adenovirus 2, Adenovirus 3, and Adenovirus 5.

The adenoviral vector, in general, also includes DNA encoding at least one therapeutic agent. The term "therapeutic" is used in a generic sense and includes treating agents, prophylactic agents, and replacement agents.

DNA sequences encoding therapeutic agents which may be placed into the adenoviral vector include, but are not limited to, the CFTR gene; genes encoding interferons such as Interferon- α , Interferon- β , and Interferon- γ ; genes encoding interleukins such as IL-1, IL-1 β , and Interleukins 2 through 14; genes encoding GM-CSF; genes encoding adenosine deaminase, or ADA; genes which encode cellular growth

3351

factors, such as lymphokines, which are growth factors for lymphocytes; genes encoding clotting factors such as Factor VIII and Factor IX; T-cell receptors; the LDL receptor, ApoB, ApoC, ApoAI and other genes involved in cholesterol transport and metabolism; the alpha-1 antitrypsin (αIAT) gene; lung surfactant protein genes, such as the SP-A, SP-B, and SP-C genes; the insulin gene, negative selective markers or "suicide" genes, such as viral thymidine kinase genes, such as the Herpes Simplex Virus thymidine kinase gene, the cytomegalovirus virus thymidine kinase gene, and the varicella-zoster virus thymidine kinase gene; Fc receptors for antigen-binding domains of antibodies, and antisense sequences which inhibit viral replication, such as antisense sequences which inhibit replication of hepatitis B or hepatitis non-A non-B virus.

The DNA sequence (or transgene) which encodes the therapeutic agent may be genomic DNA or may be a cDNA The DNA sequence also may be the native DNA sequence or an allelic variant thereof. The term "allelic variant" as used herein means that the allelic variant is an alternative form of the native DNA sequence which may have a of one or more addition substitution, deletion, or nucleotides, which does not alter substantially the function of the encoded protein or polypeptide or fragment or derivative thereof. In one embodiment, the DNA sequence may further include a leader sequence or portion thereof, a secretory signal or portion thereof and/or may further include a trailer sequence or portion thereof.

The DNA sequence encoding at least one therapeutic agent is under the control of a suitable promoter. Suitable promoters which may be employed include, but are not limited to, adenoviral promoters, such as the adenoviral major late promoter; or hetorologous promoters, such as the cytomegalovirus (CMV) promoter; the Rous Sarcoma Virus (RSV) promoter; inducible promoters, such as the MMT promoter, the

PCT/US95/15947

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metallothionein promoter; heat shock promoters; the albumin promoter; and the ApoAI promoter. It is to be understood, however, that the scope of the present invention is not to be limited to specific foreign genes or promoters.

Such vectors may be assembled by direct <u>in vitro</u> ligation from combinations of plasmids containing portions of modified or unmodified virus genome or plasmids and fragments derived directly from a linear adenoviral genome, such as the Adenovirus 5 genome (ATCC No. VR-5) or Adenovirus 5 derived viruses containing mutations or deletions as described herein.

In another alternative, the vectors can be assembled by homologous recombination, within a eukaryotic cell, between a plasmid clone containing a portion of the adenoviral genome (such as the Adenovirus 5 genome or the adenovirus 5 K3-mutant Ad dl327 (Thimmapaya, et al., Cell, Vol. 31, pg. 543 (1983)) with the desired modifications, and a second plasmid (such as, for example pAvS6, as shown in Figure 24), containing the left adenoviral ITR, an E1 region deletion, and the desired trans gene. Alternatively, homologous recombination may be carried out between a plasmid clone and a fragment derived directly from a linear adenovirus (such as Adenovirus 5, or Ad dl327 or an Adenovirus 5 derived virus containing mutations or deletions as described herein) genome.

The vector then is transfected into a cell line capable of complementing the function of any essential genes deleted from the viral vector, in order to generate infectious viral particles. The cell line in general is a cell line which is infectable and able to support adenovirus or adenoviral vector growth, provide for continued virus production in the presence of glucocorticoid hormones, and is responsive to glucocorticoid hormones (i.e., the cell line is capable of expressing a glucocorticoid hormone receptor). Cell lines which may be transfected with the essential adenoviral genes,

and thus may be employed for generating the infectious adenoviral particles include, but are not limited to, the A549, KB, and HEp-2 cell lines.

Because the expression of some viral genes may be toxic to cells, the E1 region, as well as the E2a, E2b, and/or E4 regions, may be under the control of an inducible promoter. Such inducible may include, but are not limited to, the mouse mammary tumor virus (MMTV) promoter (Archer, et al., Science, Vol. 255, pgs. 1573-1576 (March 20, 1992)); the synthetic minimal glucocorticoid response element promoter GRES (Mader, et al., Proc. Nat. Acad. Sci., Vol. 90, pgs. 5603-5607 (June 1993)); or the tetracycline-responsive promoters (Gossen, et al., Proc. Nat. Acad. Sci., Vol. 89, pgs. 5547-5551 (June 1992)). In another alternative, the E1 region is under the control of an inducible promoter, and the E2a, E2b, and/or E4 regions are under the control of their native promoters. In such alternative, the native promoters are transactivated by expression of the E1 region.

In one embodiment, the cell line includes the entire adenoviral R4 region with its native promoter region, and the R1a region or the entire R1 region (including the R1a and R1b regions) under the control of a regulatable or inducible promoter, such as, for example, the mouse mammary tumor virus (or MMTV) promoter, which is a hormone inducible promoter, or other such promoters containing glucocorticoid responsive elements (GRE's) for transcriptional control. In another embodiment, the R4 DNA sequence also is expressed from a regulatable promoter, such as the MMTV promoter. The R1 and R4 DNA sequences may be included in one expression vehicle, or may be included in separate expression vehicles. Preferably, the expression vehicles are plasmid vectors which integrate with the genome of the cell line.

In one embodiment, the producer cell line includes a first plasmid which includes the Ela or the entire El DNA sequence under the control of an inducible promoter (such as

WO 96/18418 PCT/US95/15947

the MMTV promoter), and a second plasmid Recluding the E4 DNA sequence under the control of its native promoter. an embodiment, the producer cell line is transduced with the adenoviral vector which does not include the E1 and E4 DNA sequences. The cells then are exposed to an inducing agent, (such as, for example, dexamethasone when an MMTV promoter is employed), which activates the inducible promoter, such as the MMTV promoter, thereby initiating expression of the El provides the E1 DNA expression of The DNA. transactivation of the E4 promoter, thereby providing for expression of the E4 DNA. The expression of the E1 and E4 DNA thus enables the generation of infectious adenoviral particles.

In another embodiment, the producer cell line includes a first plasmid which includes the Ela or the entire El DNA sequence under the control of an inducible promoter (such as the MMTV promoter or a promoter containing a glucocorticoid responsive element), and a second plasmid including the E4 DNA sequence under the control of an inducible promoter (such containing a promoter promoter or MMTV glucocorticoid responsive element). The producer cell line then is transduced with the adenoviral plasmid vector which does not include the B1 and B4 DNA sequences. The cells then are exposed to the inducing agent (such as, for example, dexamethasone), which activates the promoters (such as the MMTV promoters or promoters containing a glucocorticoid responsive element) controlling the E1 and E4 DNA sequences, thereby initiating expression of the E1 and E4 DNA sequences, whereby the expression of such sequences enables the generation of the infectious adenoviral particles. be understood, however, that the inducible or regulatable promoters controlling the E1 and E4 DNA sequences need not be identical.

In another embodiment, an adenoviral vector wherein the Ela or entire El DNA sequence, and the E2a DNA sequence, or

E2a and E2b DNA sequences, have been deleted, is transfect; into a cell line capable of complementing the function of t Ela or entire El, and E2a or E2a and E2b DNA sequences, as well as any other essential genes deleted from the viral vector, in order to generate infectious viral particles. one embodiment, the cell line includes the Ela or entire El region under the control of a promoter which may be its native promoter or a regulatable promoter, such as, for example, the mouse mammary tumor virus (or MMTV) promoter, or other such promoters containing glucocorticoid responsive elements (GRE's) for transcriptional control. The cell line also includes the adenoviral E2a region or E2a and E2b regions under the control of its native promoter region, or a regulatable promoter such as those hereinabove described. The Ela or entire El, and E2a or E2a and E2b DNA sequences, may be included in one expression vehicle, or may be included in separate expression vehicles.

In one embodiment, the producer cell line includes a first plasmid including the E1 DNA sequence under the control of a promoter containing a glucorticoid responsive element, and a second plasmid including the B2a DNA sequence under the control of the MMTV promoter. The producer cell line is transduced with the adenoviral plasmid vector which does not include the E1 and E2a DNA sequences. The cells then are exposed to a glucocorticoid hormone such as dexamethasone for activation of the El promoter and for activation of the MMTV promoter, thereby initiating expression of the R1 and R2a DNA of the generation infectious and enabling sequences adenoviral particles.

In another embodiment, the producer cell line includes a first plasmid including the E1 DNA sequence under the control of an inducible promoter (such as the MMTV promoter or other promoters containing a glucocorticoid responsive element), and a second plasmid including the E2a DNA sequence under the control of its native promoter. Expression of the



El and E2a DNA sequences is accomplished by the same approach as carried out with respect to the producer cell line including the E1 DNA sequence under the control of an inducible promoter, and the E4 DNA sequence under the control of its native promoter.

In yet another embodiment, the producer cell line includes a first plasmid including the E1 DNA sequence under the control of the MMTV promoter, and a second plasmid including the E2a DNA sequence under the control of the MMTV promoter. Expression of the E1 and E2a DNA sequences, leading to the generation of infectious adenoviral particles, is accomplished by exposing the cells to dexamethasone, thereby activating the MMTV promoters controlling the E1 and E2a DNA sequences.

In yet another embodiment, an adenoviral vector, wherein the E1, E2a, and E4 DNA sequences have been deleted, is transfected into a cell line which includes the adenoviral If the vector also includes a E1, E2a, and E4 regions. deletion of the B2b DNA sequence, the cell line also will include the E2b region. Each of the E1, E2a, and E4 regions may be under the control of their native promoters, or any or all of the E1, E2a, and E4 regions may be under the control of an inducible or regulatory promoter, such as those hereinabove described. The E1, E2a, and E4 DNA sequences may be included in one expression vehicle, or the E1, E2a, and E4 DNA sequences may each be included in separate expression vehicles, or two of the E1, E2a, and E4 DNA sequences may be included in one expression vehicle, and the other of the El, and E4 DNA sequences may be included in another expression vehicle. Expression of the E1, E2a and E4 DNA sequences may be accomplished by employing approaches similar to those hereinabove described.

Uses of the adenoviral vector particles of the present invention include the transduction of eukaryotic cells <u>in vivo</u> or <u>in vitro</u> as part of a gene therapy procedure, and the

PCT/US95/15947

transduction of cells <u>in vitro</u> for the <u>in vitro</u> production of desired proteins or therapeutic agents.

In one embodiment, the adenoviral vector particles are administered <u>in vivo</u> in an amount effective to provide a therapeutic effect in a host.

In one embodiment, the vector may be administered in an amount of from 1 plaque forming unit to about 10¹⁴ plaque forming units, preferably from about 10⁶ plaque forming units to about 10¹³ plaque forming units. The host may be a mammalian host, including human or non-human primate hosts.

The infectious adenoviral vector particles are administered to the lung when a disease or disorder of the lung (such as, for example, cystic fibrosis) is to be treated. Such administration may be, for example, by aerosolized inhalation or brochoscopic instillation, or via intranasal or intratracheal instillation.

In another embodiment, the infectious adenoviral vector particles are administered systemically, such as, for example, by intravenous administration (such as, for example, portal vein injection or peripheral vein injection), intramuscular administration, intraperitoneal administration, intratracheal administration.

The adenoviral vector particles may be administered in combination with a pharmaceutically acceptable carrier suitable for administration to a patient. The carrier may be a liquid carrier (for example, a saline solution), or a solid carrier, such as, for example, microcarrier beads.

Cells which may be infected by the infectious adenoviral particles include, but are not limited to, primary cells, such as primary nucleated blood cells, such as leukocytes, granulocytes, monocytes, macrophages, lymphocytes (including T-lymphocytes and B-lymphocytes), totipotent stem cells, and tumor infiltrating lymphocytes (TIL cells); bone marrow cells; endothelial cells; activated endothelial cells; epithelial cells; lung cells; keratinocytes; stem cells;

WU 90/18418 PCT/US95/15947

hepatocytes, including hepatocyte precursor cells; fibroblasts; mesenchymal cells; mesothelial cells; parenchymal cells; vascular smooth muscle cells; brain cells and other neural cells; gut enterocytes; gut stem cells; and myoblasts.

The infected cells are useful in the treatment of a variety of diseases including but not limited to adenosine deaminase deficiency, sickle cell anemia, thalassemia, hemophilia, diabetes, α -antitrypsin deficiency, brain disorders such as Alzheimer's disease, phenylketonuria and other illnesses such as growth disorders and heart diseases, for example, those caused by alterations in the way cholesterol is metabolized and defects of the immune system.

In one embodiment, the adenoviral vector particles may be used to infect lung cells, and such adenoviral vector particles may include the CFTR gene, which is useful in the treatment of cystic fibrosis. In another embodiment, the adenoviral vector may include a gene(s) encoding a lung surfactant protein, such as SP-A, SP-B, or SP-C, whereby the adenoviral vector is employed to treat lung surfactant protein deficiency states.

In another embodiment, the adenoviral vector particles may be used to infect liver cells, and such adenoviral vector particles may include gene(s) encoding clotting factor(s), such as Factor VIII and Factor IX, which are useful in the treatment of hemophilia.

In another embodiment, the adenoviral vector particles may be used to infect liver cells, and such adenoviral vector particles may include gene(s) encoding polypeptides or proteins which are useful in prevention and therapy of an acquired or an inherited defect in hepatocyte (liver) function. For example, they can be used to correct an inherited deficiency of the low density lipoprotein (LDL) receptor.

In another embodiment, the adenoviral particles may be used to infect liver cells, whereby the adenoviral particles include a gene encoding a therapeutic agent employed to treat acquired infectious diseases, such as diseases resulting from For example, the infectious adenoviral viral infection. particles may be employed to treat viral hepatitis, particularly hepatitis B or non-A non-B hepatitis. example, an infectious adenoviral particle containing a gene encoding an anti-sense gene could be employed to infect liver cells to inhibit viral replication. In this case, the infectious adenoviral particle, which includes a vector including a structural hepatitis gene in the reverse or opposite orientation, would be introduced into liver cells, resulting in production in the infected liver cells of an anti-sense gene capable of inactivating the hepatitis virus or its RNA transcripts. Alternatively, the liver cells may be infected with an infectious adenoviral particle which includes a gene which encodes a protein, such as, for example, α -interferon, which may confer resistance to the hepatitis virus.

In yet another embodiment, an adenoviral vector in accordance with the present invention may include a negative selective marker, or "suicide" gene, such as the Herpes Simplex Virus thymidine kinase (TK) gene. Such a vector may be employed in the treatment of tumors, including cancerous and non-malignant tumors, by administering the adenoviral vector to a patient, such as, for example, by direct injection of the adenoviral vector into the tumor, whereby the adenoviral vector transduces the tumor cells. After the cells are transduced with the adenoviral vector, an interaction agent, such as, for example, ganciclovir, is administered to the patient, whereby the transduced tumor cells are killed.

In another embodiment, the viral particles, which include at least one DNA sequence encoding a therapeutic

WO 96/18418 PCT/US95/15947

agent, may be administered to an animal in order to use such animal as a model for studying a disease or disorder and the treatment thereof. For example, an adenoviral vector containing a DNA sequence encoding a therapeutic agent may be given to an animal which is deficient in such therapeutic agent. Subsequent to the administration of such vector containing the DNA sequence encoding the therapeutic agent, the animal is evaluated for expression of such therapeutic agent. From the results of such a study, one then may determine how such adenoviral vectors may be administered to human patients for the treatment of the disease or disorder associated with the deficiency of the therapeutic agent.

In another embodiment, the adenoviral vector particles may be employed to infect eukaryotic cells <u>in vitro</u>. The eukaryotic cells may be those as hereinabove described. Such eukaryotic cells then may be administered to a host as part of a gene therapy procedure in amounts effective to produce a therapeutic effect in a host.

Alternatively, the vector particles include a gene encoding a desired protein or therapeutic agent may be employed to infect a desired cell line in vitro, whereby the infected cells produce a desired protein or therapeutic agent in vitro.

EXAMPLES

The invention now will be described with respect to the following examples; however, the scope of the present invention is not intended to be limited thereby.

WÖ 96/18418 PCT/US95/15947

Example 1

Development of Cell Lines for Complementation of E1'/E2a' or E1'/E4' Vectors

Construction of plasmid having Adenovirus 5 El sequence.

The Adenovirus 5 genome was digested with ScaI enzyme, separated on an agarose gel, and the 6,095 bp fragment comprising the left end of the virus genome was isolated. The complete Adenovirus 5 genome is registered as Genbank accession #M73260, incorporated herein by reference, and the virus is available from the American Type Culture Collection, Rockville, Maryland, U.S.A., under accession number VR-5. The ScaI 6,095 bp fragment was digested further with ClaI at bp 917 and BglII at bp 3,328. The resulting 2,411 bp ClaI to BglII fragment was purified from an agarose gel and ligated into the superlinker shuttle plasmid pSE280 (Invitrogen, San Diego, CA), which was digested with ClaI and BglII, to form pSE280-E. (Figure 1).

Polymerase chain reaction (PCR) was performed to synthesize DNA encoding an XhoI and SalI restriction site contiguous with Adenovirus 5 DNA bp 552 through 924. The primers which were employed were as follows:

5' end, Ad5 bp 552-585:

(%)

- 5'-GTCACTCGAGGACTCGGTC-GACTGAAAATGAGACATATTATCTGCCACGGACC-3'
- 3' end, Ad5 bp 922-891:
- 5'-CGAGATCGATCACCTCCGGTACAAGGTTTGGCATAG-3'

This amplified DNA fragment (sometimes hereinafter referred to as Fragment A) then was digested with XhoI and ClaI, which cleaves at the native ClaI site (bp 917), and ligated to the XhoI and ClaI sites of pSE280-E, thus reconstituting the 5' end of the E1 region beginning 8 bp upstream of the ATG codon.

PCR then was performed to amplify Adenovirus 5 DNA from bp 3,323 through 4,090 contiguous with an EcoRI restriction site. The primers which were employed were as follows:

5' end, Ad5 bp 3323-3360:

WO 90/18418 PCT/US95/15947

5'-CATGAAGATCTGGAAGGTGCTGAGGTACGATGAGACC

3' end, Ad5 bp 4090-4060:

A : 1

5'-GCGACTTAAGCAGTCAGCTG-AGACAGCAAGACACTTGCTTGATCCAAATCC-3'

This amplified DNA fragment (sometimes hereinafter referred to as Fragment B) was digested with BglII, thereby cutting at the Adenovirus 5 BglII site (bp 3,382) and EcoRI, and ligated to the BglII and EcoRI sites of pSE280-AE to reconstruct the complete Ela and Elb region from Adenovirus 5 bp 552 through 4,090. The resulting plasmid is referred to as pSE280-E1 (Figure 1).

The intact Ela/Elb region was excised from pSE280-El by cutting with EcoRI, blunting with Klenow and cutting with SalI. The 3,567 bp fragment was purified from an agarose gel and ligated to the expression plasmid pMAMneo (Clonetech, Palo Alto, CA), which was prepared similarly by cutting with KhoI at bp 3,405 of pMAMneo Luc (Figure 2), blunting with Klenow, and cutting with SalI at bp 1,465. The 8,399 bp fragment was gel purified, phosphatased, and ligated to the Ela/Elb fragment hereinabove described to form pMAMneo-El. (Figure 3.)

Bacterial transformants containing the final pMAMneo-El construct were identified. Plasmid DNA was prepared and purified by banding in CsTFA. Circular plasmid DNA was linearized at the XmnI site within the ampicillin resistance gene of pMAMneo-El, and purified further by phenol/chloroform extraction and ethanol precipitation prior to use for transfection of cells.

An alternative construction containing the intact Ela/b region under the control of the synthetic promoter GRE5 was prepared as follows. The intact Ela/b region was excised from pSE280-El, which was modified previously to contain a BamHI site 3' to the El gene, by digesting with XhoI and BamHI. The XhoI to BamHI fragment containing the Ela/b fragment was cloned into the unique XhoI and BamHI sites of

pGRE5-2/EBV (Figure 4, U.S. Biochemicals, Cleveland, Ohio) to form pGRE5-E1 (Figure 5).

Construction of plasmid including Adenovirus 5 B2a sequence.

The Adenovirus 5 genome was digested with BamHI and SpeI, which cut at bp 21,562 and 27,080, respectively. Fragments were separated on an agarose gel and the 5,518 bp BamHI to SpeI fragment was isolated. The 5,518 bp BamHI to SpeI fragment was digested further with SmaI, which cuts at bp 23,912. The resulting 2,350 bp BamHI to SmaI fragment was purified from an agarose gel, and ligated into the superlinker shuttle plasmid pSE280, and digested with BamHI and SmaI to form pSE280-E2 BamHI-SmaI (Figure 6).

PCR then was performed to amplify Adenovirus 5 DNA from the SmaI site at bp 23,912 through 24,730 contiguous with NheI and EcoRI restriction sites. The primers which were employed were as follows:

- 5' end, Ad5 bp 24,732-24,708:
- 5'-CACGAATTCGTCAGCGCTTCTCGTCGCGTCCAAGACCC-3'
- 3' end, Ad5 bp 23,912-23,934:

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5'-CACCCCGGGAGGCGGCGACGGGACGGG-3'

This amplified DNA fragment was digested with SmaI and EcoRI, and ligated to the SmaI and EcoRI sites of pSE280-E2 Bam-Sma to reconstruct the complete E2a region from Ad5 bp 24,730 through 21,562. The resulting construct is pSE280-E2a. (Figure 6.)

In order to convert the BamHI site at the 3' end of E2a to a SalI site, the E2a region was excised from pSE280-E2a by cutting with BamHI and NheI, and recloned into the unique BamHI and NheI sites of pSE280. (Figure 6.) Subsequently, the E2a region was excised from this construction with NheI and SalI in order to clone into the NheI and SalI sites of the pMAMneo multiple cloning site in a 5' to 3' orientation, respectively. The resulting construct is pMAMneo E2a. (Figure 6).

WO 96/18418 PCT/US95/15947

Bacterial transformants containing the final pMAMneo-E2a were identified. Plasmid DNA was prepared and purified by banding in CsTFA. Circular plasmid DNA was linearized at the XmnI site within the ampicillin resistance gene of pMAMneo-E2a, and further purified by the phenol/chloroform extraction and ethanol precipitation prior to use for transfection of cells. Alternatively, the plasmid was linearized with BclII, which cuts in the reading frame of the neo^R gene, thereby inactivating it. This form of the plasmid was used where cotransfection with an alternative selectable marker was required.

Construction of plasmid including Adenovirus 5 ITR/E4 region.

The Adenovirus 5 genome was digested with SpeI, which cuts at bp 27,082, and the two fragments separated on an agarose gel. The 8,853 bp fragment comprising the right end of Adenovirus 5 was isolated.

The 8,853 bp right end fragment was digested further with StuI at bp 31,956 and Eco47-III at bp 35,503. Fragments were separated on an agarose gel and the 3,547 bp StuI to Eco47-III fragment was isolated. This fragment was ligated to the superlinker shuttle plasmid pSE380 (Invitrogen) (Figure 7), digested with StuI and Eco47-III to construct pSE380-E4 (Figure 8).

PCR was performed to amplify Adenovirus 5 DNA from bp 35,499 through the right ITR ending at bp 35,935, and contiguous with a BamHI restriction site. The PCR primers which were employed were as follows:

5' end, Ad5 bp 35,935-35,908:

ad

- 5'-GTGGGATCCATCATCAATAATATACCTTATTTTGGA-3'
- 3' end, Ad5 bp 35,503-35,536:
- 5'-ATACAGCGCTTCCACAGCGGCAGCCATAACAGTC-3'

This amplified DNA was digested with BamHI and Eco47-III, which cuts at the Adenovirus 5 Eco47-III site bp 35,503, and ligated to the unique BamHI and Eco47-III sites of pSE380-E4 to form pSE380-ITR/E4. (Figure 8.)

Construction of a plasmid containing the E4 region under the control of an inducible promoter.

A plasmid expressing all of the adenovirus 5 E4 open reading frames under the control of the glucocorticoid inducible GRE5 promoter, and containing a puromycin resistance gene under the control of the control of the SV40 promoter was constructed as follows.

The construct pSE380-E4 (Figure 8) was cut with Eco47-3 and BamHI and ligated to a synthetic oligonucleotide of 33 bp, which includes the Eco47-3 site at Adenovirus 5 bp 35,503 through the E4 ORF1 ATG at bp 35,522. The resulting construct, pSE380E4-ATG (Figure 9) contains all E4 open reading frames without a promoter. The E4 ORF's were excised from pSE380E4/ATG as an XhoI to BamHI fragment and ligated to the XhoI and BglII sites within the multiple cloning site of the expression shuttle plasmid pSE380GRE5/SV40Puro^R, to generate pGRE5/E4Puro (Figure 9). The expression shuttle plasmid pSE380GRE5/SV40Puro^R was constructed from pSE380 (Figure 7), the GRE5 promoter excised from pGRE5/EBV (Figure 4), and a puromycin resistance gene.

Transfection and selection of cells.

general, this process involved the sequential introduction, by calcium phosphate precipitation, or other means of DNA delivery, of two plasmid constructions each with a different viral gene, into a single tissue culture cell. The cells were transfected with a first construct and selected for expression of the associated drug resistance gene to establish stable integrants. Individual cell clones were established and assayed for function of the introduced Appropriate candidate clones then were viral gene. transfected with a second construct including a second viral gene and a second selectable marker. Transfected cells then were selected to establish stable integrants of the second construct, and cell clones were established. Cell clones were assayed for functional expression of both viral genes.

PCT/US95/15947

WU 90/18418

297

In order to determine the most suitable cell lines for the above-mentioned transfections, sequential transfections and selections were carried out with the following parental cell types:

A549 (ATCC Accession No. CCL-185);

Hep-2 (ATCC Accession No. CCL-23); or

KB (ATCC Accession No. CCL-17).

Appropriate selection conditions were established for both G418 and hygromycin B for all three cell lines by standard kill curve determination.

Transfection of cell lines with plasmids including E1 and E2a regions.

pMAMneo-B2a was digested with BclII to linearize the plasmid within the neoR reading frame and inactivate the gene. The cell lines hereinabove described were co-transfected with the linear pMAMneoR-E2a and a selectable marker plasmid consisting of a eukaryotic expression cassette containing an hygromycin resistance gene, SV40 promoter, a polyadenylation signal cloned into the multiple cloning site of a pTZ18R (Figure 10), at a 10:1 molar ratio. 48 hours after transfection, the cells were placed under hygromycin selection and maintained in selection until drug resistant colonies arose. The clones were isolated and screened for E2a expression by staining for E2a protein with a polyclonal and visualizing by immunofluorescence. function was screened by complementation of the temperaturesensitive mutant Ad5ts125 virus which contains a temperaturesensitive mutation in the E2a gene. (Van Der Vliet, et al., J. Virology, Vol. 15, pgs. 348-354 (1975)). Positive clones were identified and transfected with pMAMneoEl linearized with XmnI within the ampicillin resistance gene. The cells were selected for G418 resistance. G418-resistant colonies were screened for B1 expression by staining with a monoclonal antibody for the El protein (Oncogene Sciences, Uniondale, N.Y.). El function was assayed by ability to complement an WU 96/18416 PC1/US95/15947

E1-deleted vector such as pAvS6 (Figure 20) described hereinbelow.

In a preferred alternative, pMAMNeo-E2a was linearized with XmmI with the Amp^R gene, introduced into cells by transfection, and cells were selected for stable integration of this plasmid by G418 selection. Clones expressing the E2a gene were identified and used for transfection with the 7 kb EcoRV to XmmI fragment from pGRE5-E1 (Figure 5), which contains the GRE5 promoted E1a/b region plus the hygromycin^R gene. Cells were selected for hygromycin resistance and assayed for both E1a/b and E2a expression.

Transfection of cell lines with plasmids including E1 and E4 regions.

Circular plasmid pSE380-E4 was co-transfected into the cell lines hereinabove described with a hygromycin resistance plasmid as described above at a 10:1 molar ratio, 48 hours after transfection, the cells were placed under hygromycin selection and maintained in selection until drug resistant colonies arose. The clones then were isolated and screened Specifically, clones were tested for their for E4 function. ability to complement the Adenovirus 5 mutant dl 1011 (Bridge, et al. <u>Virology</u>, Vol. 193, pgs. 794-801 (1993)), which contains a deletion of all E4 open reading frames, and to produce infectious virus of the K4 genotype. clones were identified and transfected with pMAMneoE1 linearized with XmmI within the ampicillin resistance gene. The cells then were selected for G418 resistance. G418resistant colonies then were screened for El function.

4555

Transfection of cells containing the inducible B1 and inducible B2a plasmids with an inducible B4 expression plasmid.

In the preferred alternative described above for the generation of cells containing pMAM-neo/E2a and pGRE5-E1, clones were identified that inducibly express both viral genes. These cell lines are modified further by transfection

PCT/US95/15947

with the pGRE5/E4/SV40Puro' plasmid (Fig. 9) and selection for puromycin resistant colonies. Cell clones are selected, expanded, and analyzed for inducible E4 function by assaying for dexamethasone-dependent complementation of the E4 deleted virus Ad5dl1011. (Bridge, et al., Virology, Vol. 193, pgs. 794-801 (1993)). In another alternative, the E4 gene would be operationally linked to a tetracycline-inducible promoters.

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250

Example 2

Development of Vectors with Deletions of E1/E2a or E1/E4

Intermediates for construction of a vector with E2A deletion

Subcloning of Adenovirus 5 E2 region.

Adenovirus 5 genomic DNA was cut with BamHI (bp 21,562) and SpeI (bp 27,082) restriction enzymes, and the 5,520 bp BamHI to SpeI fragment was isolated from an agarose gel. The 5,520 bp BamHI to SpeI fragment was ligated to the pTZ18R plasmid (Pharmacia, Piscataway, NJ) (Figure 10), which was modified previously to contain an SpeI site at bp 186, to generate plasmid pTZE2A. (Figure 11.)

Deletion of the E2A open reading frame.

pTZE2A was cut with BamHI (bp 8,391) and SmaI (bp 2,350), restriction enzymes, corresponding to bp 21,562 and 23,912 of Adenovirus 5. The 6,041 bp and 2,350 bp BamHI/SmaI fragments were isolated from an agarose gel. The 2,350 bp BamHI/SmaI fragment was digested with DraI restriction enzyme (bp 22,442 of Adenovirus 5), and the 880 bp DraI to BamHI fragment was isolated from an agarose gel. The 880 bp DraI/BamHI fragment was ligated to the 6,041 bp BamHI/SmaI fragment via a cohesive BamHI to BamHI and blunt ended SmaI to DraI ligation, thereby generating plasmid pTZdlE2A. (Figure 11.) Such plasmid includes the Adenovirus 5 E2A region in which 1,470 bp of the E2A region, corresponding from the DraI site (bp 22,442) to the SmaI site (bp 23,912) have been deleted.



Subcloning of the Adenovirus 5 E3 region.

The Adenovirus 5 genome was digested with SpeI (at bp 27,082). The 8,853 bp fragment comprising the right end of Adenovirus 5 was isolated. The 8,853 bp right end fragment of Adenovirus 5 was digested with StuI (bp 31,956). The 4,874 bp SpeI/StuI fragment was isolated. This fragment was ligated to the plasmid pSE380 (Invitrogen) which was digested with StuI and SpeI, in order to generate pSE380-E3+. (Figure 12.)

Combining the E3 and E4 regions.

The plasmid SE380-E3+ was digested with SpeI and StuI, and the 4,874 bp fragment was ligated to the pSE380-ITR/E4 plasmid (Figure 8), which was digested with SpeI and StuI, to generate pSE380-E3+E4+. (Figure 13.)

The E3+/E4+ fragment then was excised as an 8,943 bp SpeI to BamHI fragment from pSE380-E3+E4+, and recloned into the SpeI and BamHI sites of pTZ18R, previously modified to remove the XbaI site, to generate pTZE3+E4+. (Figure 13.)

Deletion within the E3 region.

The plasmid pTZE3+E4+ was digested with XbaI, in order to delete 1,878 bp of the E3 region between the XbaI sites at bp 28,592 and 30,470 of the Adenovirus 5 genome, to generate pTZE3E4+. (Figure 14.)

The plasmid pTZE3E4+ was digested with SpeI and BamHI and ligated into the SpeI and BamHI sites of the pSE280 plasmid to form pSE280-E3E4+. The pSE280-E3E4+ plasmid was digested with BamHI, blunted, and recircularized to remove the BamHI site. The E3E4+ containing fragment was removed by restriction digestion with SpeI and SalI and ligated to the pSPORT-1 (Figure 15) (Gibco/BRL, Gaithersburg, MD) plasmid digested with SpeI and SalI, to generate pSPORT1E3E4+. (Figure 14.)

Combining the RAL rection with the LIRA+ rection

The plasmid psport-lessed was disposed with spel and BamHI, and ligated to the BamHI/Spel fragment from plasmid pTZdlE2A, to generate psport-lesars). (Figure 16.)

Construction containing an E2A region deligion and a modified R3 region

In another alternative, the pSPORT1/E2A-E3+E4+ (Figure 16) may be modified by inserting an XbaI fragment containing a heterologous promotor, such as SV40, operationally linked to all or a part of the E3 region open reading frames into the unique XbaI site to generate pSPORT1/E2A-E3+*E4+.

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Incorporation of R2AE3E4+ regions into a circular, non-infectious Adenovirus 5 plasmid.

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Plasmid pSPORT-1E2AE3E4+ was digested with BamHI and SalI. The fragment containing the E2AE3E4+ sequences was ligated to plasmid pBR322 (Figure 17) (Life Technology, Gaithersburg, MD) digested with BamHI and SalI, to generate pBR322E2AE3E4+. (Figure 18.)

The Adenovirus 5 genome was digested with ClaI (bp 917) and BamHI (bp 21,562), and the 20,645 bp fragment was ligated to the plasmid pBR322E2AE3E4+ (which also was digested with ClaI and BamHI), to generate pBRAd5E2AE3E4+. (Figure 18.)

Intermediates for construction of a vector with an E4 deletion.

The plasmid pSE380-ITR/E4 was digested with BpUll02 (bp 33,129) and Eco47-3 (bp 35,503) to remove all E4 open reading frames, to generate pSE380-E4. (Figure 19.)

The E4 region (StuI to BamHI) of plasmid pSE380-E3+E4+ was removed and replaced with the StuI to BamHI fragment from pSE380-E4, in order to incorporate the E4 deletion and generate pSE380-E3+E4. (Figure 19.)

In a preferred alternative, a larger E4 region deletion is made which reduces a 60 amino acid reading frame initiated

WO 96/18418 PCT/US95/15947 .

from the E4 ORF1 ATG of pSE380E4 to seven amino acids. This preferred modification, referred to as pSE380E4 (2) is constructed as shown in Figure 20. pSE380E4 is cut with Bpu 1102 (Adenovirus 5 bp 33,129) and AflII (Adenovirus 5 bp 33,104), blunt ended by filing in 5' overhangs with Klenow, and religating to form pSE380E4 (2). This alternative E4 region deletion then is excised as a StuI to BamHI fragment and substituted for the StuI to BamHI fragment in pSE380E3 *E4* (Figure 13) to generate pSE380E3 *E4* (2), as shown in Figure 20.

As a further modification, the XbaI region between Adenovirus 5 bp 28,592 and 30,470 is deleted from pSE380E3*E4*(2) in order to generate pSE380E3*E4*(2), as shown in Figure 21.

In another alternative, the XbaI region between Ad5 bp 28,592 and 30,470 in pSE380E3+E4(2); Figure 21, may be replaced with an XbaI fragment containing a heterologous promotor, such as SV40, operationally linked at some or all of the E3 region open reading frames, to create pSE380E3+*E4-(2).

Combining deletions within the B2A, R3, and R4 regions into a single plasmid

An Adenovirus 5 DNA fragment containing both the E3 region deletion and E4 region deletion in tandem is excised from pSE380E3E4'(2) by digestion with SpeI (Adenovirus 5 bp 27,082) and SalI (Figure 22). This fragment is substituted for the unique SpeI to SalI fragment in pSPORT1/E2AE3E4' (Figure 16) to form pSPORT1/E2AE3E4'(2), as shown in Figure 22.

In another alternative, the unique SpeI to SalI fragment in pSPORT1/E2A-E3-E4+ (Figure 16) may be replaced with the SpeI to SalI fragment excised from pSE380E3+*E4-(2) to create pSPORT1/E2A-E3+*E4-(2).

Virus Vector Production

Production of a vector with the genotype E1+E2AE3E4+.

WO 96/18418 PCT/US95/15947

The plasmid pSPORT-1E2AE3E4+ is digested with BamHI and SalI, and the fragment containing the E2AE3E4+ sequences is purified from an agarose gel. The unique BamHI site corresponds to the BamHI site at bp 21,562 of Adenovirus 5. The purified fragment is ligated in vitro to the left side BamHI fragment of the Adenovirus 5 genome (21,562 bp). The ligation mixture then is transfected into an appropriate cell line capable of complementing the vector E2A deletion for production of an E1+E2E3E4+ Adenovirus 5 vector. (Figure 23.)

Production of a Vector with the genotype E1+E2A-E3+*E4+

In a manner analogous to that described above and shown in Figure 23, the BamHI to SalI fragment may be isolated from psport1/E2A-E3+*E4+ and used in the ligation with the Ad5 21,562 bp left end BamHI fragment to generate an Ad5 virus with the genotype E1+E2A-E3+*E4.

Production of a vector with the genotype E1E2AE3E4+.

303

The DNA of the B1+B2B3B4+ Adenovirus 5 vector is purified and digested with ClaI. This restriction enzyme cuts the Adenovirus 5 genome at bp 917. The right side fragment is purified and co-transfected with pAvS6 (including a reporter gene or a trans-gene) into a double complementing cell line as capable of providing both E1 and E2A functions pAvS6 (Figure 24) is a shuttle plasmid which includes an adenoviral 5' ITR, and adenoviral encapsidation signal, an Ela enhancer sequence; a promoter; a tripartite leader sequence, a multiple cloning site for the insertion of foreign genes; a poly A signal; and a DNA segment corresponding to the Adenovirus 5 genome which is no longer than from base 3329 to base 6246 of the genome. in PCT Application published described further 27, 1994. Homologous October published WO94/23582, recombination between the virus DNA fragment and pAvS6 results in a linear virus genome with the genotype E1 (trans gene) E2aE3E4+. (Figure 25.) This vector may be plaque WO 96/18418 PCT/US95/15947

purified further using a double complementing cell line, E1+/E2a+.

In another alternative, a vector with the genotype E1E2A E3E4+ may be generated by ligating the left ITR through BamHI fragment of a pre-existing E1 vector to the BamHI/SalI fragment of pSPORT-1E2AE3E4 (Figure 26) to generate directly an E1 (reporter or trans-gene) E2AE3E4+ virus which will replicate when transfected into an appropriate complementing cell line. This approach, however, requires that the reporter or trans-gene does not contain a BamHI site.

In yet another alternative (Figure 27), plasmid pBRAd5-E2AE3E4+ is co-transfected with pAvS6 (containing a reporter gene or a trans-gene), into a double complementing cell line, to generate by homologous recombination an E1 (reporter or trans-gene) E2AE3E4+ vector. The vector is plaque purified using a double complementing cell line.

As an example of the practice of these methods, a vector with the genotype R1E2aE3E4⁺ was generated using the adenoviral plasmid vector pAvS6nLacZ (described in PCT application No. W095/09654), which contains a nuclear localizing beta-galactosidase cDNA. The resulting vector, Av3nLacZ (Figure 28), was generated by homologous recombination as outlined in Figure 25.

In another alternative, the vector Av3nLacZ can be used as a starting material for the production of other vectors. Av3nLacZ contains two ClaI sites, one within the lacZ coding 3' end of <u>lacZ</u> within the the region and one at polyadenylation signal. (Figure 28). By digesting Av3nLacZ with ClaI and recombining with pAvS6/transgene as described in Figure 25, other Av3-type vectors are made. This approach provides advantages in that no E1 + vector backbone is present in the recombination, and new recombinant vectors can be selected as B-gal negative when X-gal is included in the agar overlay during plaque purification.

W O 96/18418

PCT/US95/15947

Production of a vector with the genotype E1-E2A-E3+*E4+

In a manner analogous to that described above and shown in Figure 25, DNA from the virus E1+E2A-E3+*E4+ is digested with ClaI and the right side fragment used to co-transfect with pAVS6/transgene into E1/E2A expressing cells to generate, by homologous recombination, a vector with the genotype E1-E2A-E3+*E4+.

In another alternative, and in a manner analogous to that shown in Figure 26, the left end BamHI fragment from a pre-existing E1- vector may be ligated directly to the BamHI/SalI fragment from a pSPORT1/E2A-E3+*E4+ and transfected into E1/E2A expressing cells to generate a vector with the genotype E1-, transgene, E2A-E3+*E4+.

Production of a vector with the genotype E1+E4.

A purified Adenovirus 5 fragment from bp 1 to the SpeI site (bp 27,082), containing terminal protein on the 5' end, was ligated to the SpeI/BamHI fragment purified from plasmid pSE380-E3+E4' containing the E4 region deletion. The *in vitro* ligation mixture is transfected into an E4+ cell line to produce an Adenovirus 5 E1+E4' vector. (Figure 29.)

Alternatively, pSE380E3*E4*(2), as shown in Figure 20, is used to generate an E1*E3*E4*(2) virus with the E4 deletion in the same manner as outlined in Figure 29. In another alternative, pSE380E3*E4*(2), as described in Figure 21, is used in the same manner to generate an E1*E3*E4*(2) adenovirus.

Production of a vector with genotype E1 E4.

The DNA of the Adenovirus 5 E1+E4 genome is purified and digested with ClaI restriction enzyme. The right side and co-transfected with pAvS6 purified is fragment (containing a reporter or trans-gene) into a complementing cell line expressing E1 and E4. recombination between the E4 viral fragment and pAvS6 will result in a vector with the genotype El (reporter or transgene) E2A+E3+E4:. The vector (Figure 30) is plaque purified using a double complementing cell line.

WO 96/18418

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In another alternative, the SpeI/BamHI fragme SE380-E3+E4 is ligated directly to the left end frag a pre-existing E1 vector digested with SpeI to generat (reporter or trans-gene)) E4 vector. (Figure 31.) approach, however, requires that the reporter or tran does not contain an SpeI site.

In other alternatives, by using either the homologous recombination or the <u>in vitro</u> ligation approaches described above (Figures 30 and 31), and the E1⁺E3⁺E4⁻(2) or E1⁺E3⁻E4⁻(2) viruses hereinabove described, transgene-containing vectors of the types E1⁻(transgene) E3⁺E4⁻(2) or E1⁻(transgene) E3⁻E4⁻(2) are generated.

Example 3

Production of an Adenovirus vector with the genotype E1', E2A', E3', E4'.

Adenovirus vectors with deletions in 3 essential genes (E1, E2A, E4), referred to as triple deletion vectors or AV4 generation vectors are constructed as follows. Using Av3nLacZ as a starting material, AV4nLacZ is generated by either homologous recombination or in vitro ligation. Using AV4nLacZ as a starting material, any other transgene containing Av4 type vector can be constructed.

In one alternative, the unique BamHI to SalI fragment from pSPORT1/E2AE3E4'(2) is isolated and co-transfected with the left end fragment of Av3nLacZ cut at the unique SrfI restriction site (bp 27,098 of AV3nLacZ (Figs. 28, 32) into a triple complementing cell line (as described in Example 1 hereinabove) expressing E1, E2A, and E4 viral genes. Homologous recombination between the these two DNA fragments in the overlapping region between BamHI and SrfI will result in a vector with the genotype E1, nLacZ, E2A', E3', E4' (Fig. 32).

In another alternative, the unique SrfI to SalI fragment from pSPORT1/E2AE3E4 (2) may be isolated and directly ligated to the left end fragment of AV3nLacZ cut with SrfI at bp

WO 96/18418 PCT/US95/15947

27,098 to yield a vector with the genotype E1, nLacZ, E2A, E3, E4. This ligation mixture may then be transfected into the triple complementing cell line (E1⁺, E2A⁺, E4⁺) to produce AV4nLacZ (Fig. 33).

In another alternative, AV4 vectors containing transgenes other than nLacZ are constructed. Purified DNA from AV4nLacZ is digested with ClaI (See Fig. 34), and the right end viral fragment is purified. This fragment is cotransfected with the desired pAVS6/transgene construct into triple complementing cells (E1+, E2A+, E4+) and by homologous recombination to yield an AV4 type vector with the genotype E1 , transgene, E2A, E3, E4 (Fig. 34).

Production of an Adenovirus vector with the genotype R1-R2A-R3+*R4-

In one alternative, and in a manner analogous to the process shown in Figure 32, a unique BamHI/SalI fragment from pSPORT1/E2A-E3+*E4-(2) may be isolated and co-transfected with the left end fragment of AV3nLacZ cut at the unique SrfI restriction site into cells expressing E1, E2A, and E4 in order to generate, by homologous recombination, a vector with the genotype E1-,nLacZ,E2A-E3+*E4(2) or AV4nLacZE3+*.

In another alternative, and in a manner analogous to the process shown in Figure 33, the unique SrfI to SalI fragment from pSPORT1/B2A-B3+*B4-(2) may be isolated and used to ligate directly to the left end fragment of AV3nLacZ cut with SrfI. The ligation product is then transfected into cells expressing B1, B2A, and B4 to generate a vector with the genotype B1-,nLacZ,E2A-B3+*B4-(2), or AV4nLacZB3+*.

In another alternative, AV4 vectors containing a transgene other than nLacZ and containing the modified E3+* region can be generated in a fashion analogous to that shown in Figure 34. AV4nLacZE3+* is cut with ClaI and cotransfected with pAV6/transgene into cells expressing E1, E2A, and E4 to generate a vector with the genotype E1-, transgene, E2A-E3+*E4-(2).

WO 96/18418 PCT/US95/15947

Example 4

Development of Recombinant Adenovirus Vectors
with Deletions of E1/E2a, or E1/E4 or E1/E2a/E4
which express the normal human CFTR cDNA.

pAvS6 (Figure 24) was linearized with EcoRV. The normal human CFTR cDNA sequence (nucleotides 75 to 4,725; for numbering see Gen Bank accession number M28668) was removed from plasmid pBQ4-7 (Figure 35) by PstI digestion followed by blunting of the CFTR cDNA ends with T4 polymerase (pBQ4-7 was provided by L.-C.Tsui, the Hospital for Sick Children, Toronto, Canada.) This CFTR cDNA was then inserted into the linearized pAvS6 plasmid so as to create an operational linkage between the RSV promoter and the 5' end of the coding sequence of the CFTR cDNA. The resulting plasmid pAvS6 CFTR (Figure 35) was linearized by digestion with KpnI and cotransfected into E1/E2a, E1/E4 or E1/E2a/E4 expressing cells along with the large Cla1 fragment of Av3LacZ for E1/E2, Av5LacZ for E1/E2/E4, or the large ClaI fragment from Av4nLacZ, respectively, as described above.

iE1/E2a cells cotransfected as described above were then overlaid with agent and cultured in a humidified atmosphere containing 5% CO₂ at 37°C until formation of virus plaques. Plaques were picked and the recombinant adenovirus vectors further plaque-purified, amplified and titered as previously described (Rosenfeld, et al., Cell, Vol. 68, pgs. 143-155 (1992). Adenoviral vectors are evaluated for deletion of Ad genes E1/E2a, E1/E4 or E1/E2/E4 and inclusion of part or all of the normal human CFTR cDNA as previously described (Mittereder, et al., Human Gene Therapy, Vol. 5, pgs. 717-729 (1994). The ability of such CFTR expressing adenovirus vectors to correct the Cl secretory defect of CF epithelial cell lines was tested as previously described. (Mittereder, et al., 1994).

Example 5

Demonstration that E1/E2a-deleted vectors have reduced

PCT/US95/15947

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expression of adenoviral late gene products.

This example demonstrates that third generation vectors which harbor deletions in all or part of E1/E2a with or without deletions of E3 region sequences have reduced potential for elicitation of a host immune response. The expression of hexon gene was measured by metabolic labeling. Further, this expression was compared among a first generation vector (e.g., E1/E3 deleted), a second generation vector (e.g., E1 deleted containing a temperature sensitive mutation in the E2a gene (Englehardt, et al., Proc. Nat. Acad. Sci., Vol. 91, pgs. 6196-6200 (1994)) and the third generation vector described above.

To evaluate the performance of these vectors with regard to the decrease in late gene expression due to B2a deletion, A549 cells were infected with Av1LacZ4 (50-500 iu/cell) Av2Luc1, which is an B1 deleted vector containing a temperature sensitive mutation in the E2a gene, (50-500 iu/cell). or Av3LacZ1 (500-3000 luciferase gene. in/cell) and cultured in a humidified atmosphere with 5% Co₂ at 37°C. As controls, either iE1/E2a cells or 293 cells (expressing the inducible R1 and R2a regions of Ad5) were infected with AvlLacZ4, Av2Luc1 or Av3LacZ1 at an approximate multiplicity of infection (M.O.I.) of 10 infectious units Media was then changed to methionine-free DMEM containing 35S-Met (Mittereder, et al., 1994) cultured for an additional 24 hours. Metabolically labelled cells were then washed in phosphate buffered saline, lysed and then scraped into 800 μl of Ripa buffer containing antiproteases (PMSF, leupeptide and proteinase). thorough mixing to homogenize the cells, 35S-Met incorporation by quantified was labeled protein tricholoracetate precipitation. Aliquots of each incubation containing 3x106 cpm were immunoprecipitated using antiadenovirus 5 hexon antibody and evaluated using SDS-page as previously described (Mittereder, et al., Human Gene Therapy,

WO 96/18418 PCT/US95/15947

5:717-729, 1994). The results show that, at 37°C, normal body temperature, Av3LacZ1 demonstrated markedly reduced hexon gene expression in A549 cells even at a high MOI (Figure 36) and 293 cells (Figure 37, lanes 6-8) than did either Av1LacZ4 or Av2Luc1. As a control, all three vectors express similar amounts of hexon in E1/E2a expressing cells which complement the various adenoviral gene deletions within the vector (Figure 36, lanes 1-3). This demonstrates that the third generation vectors have improved (decreased) expression of adenoviral backbone genes.

Example 6

Demonstration that E1/E2a-deleted vectors have reduced replication of the adenoviral DNA genome.

In order to demonstrate that third generation (E1'/E2a') vectors with or without deletions of the E3 region sequences have improved characteristics with respect to residual viral DNA replication, metabolic labeling of vector infected cells with 32P was conducted. These experiments were carried out in a manner analogous to that described in Example 5 above. evaluate the replication defect of these vectors, 293 and iE1/E2 cells were infected with Av1LacZ4, Av2Luc1 or Av3LacZ1 at approximately 10 infectious units per cell and cultured as PO₄-free media containing ¹²P-orthophosphate and above. cultured for an additional 8 hours. Cells were then washed in phosphate buffered saline then lysed in 0.8 ml of Hirt Viral DNA was purified, digested with XbaI solution. restriction enzyme and subjected to a agarose electrophoresis and autoradiography as previously described. (Mittereder, et al., 1994).

The result demonstrate the Av3LacZ1 undergoes significantly less replication than Av2Luc1 or Av1Luc1 (Figure 38). Thus, third generation adenoviral vectors have improved function in relation to adenovirus vector DNA replication.

The disclosure of all patents, publications, (including published patent applications), and database accession numbers and depository accession numbers referenced in this specification are specifically incorporated herein by reference in their entirety to the same extent as if each such individual patent, publication, and database accession number, and depository accession number were specifically and individually indicated to be incorporated by reference.

It is to be understood, however, that the scope of the present invention is not to be limited to the specific embodiments described above. The invention may be practiced other than as particularly described and still be within the scope of the accompanying claims.



WHAT IS CLAIMED IS:

- 1. An adenoviral vector wherein the adenoviral genome has been modified to reduce the host immune and inflammatory responses to said vector.
- 2. The adenoviral vector of Claim 1 wherein said vector includes an adenoviral 5' ITR; an adenoviral 3' ITR; an adenoviral encapsidation signal; at least one DNA sequence encoding a protein or polypeptide of interest; and a promoter controlling the DNA sequence(s) encoding a protein(s) or polypeptide(s) of interest, said adenoviral vector being free of all or a portion of the adenoviral E1 and free of all or a portion of the E4 DNA sequences.
- 3. The vector of Claim 2 wherein said vector also is free of the adenoviral Ela and Elb DNA sequence.
- 4. The vector of Claim 2 wherein said vector is free of <u>at</u>
 least open reading frame 6 of the E4 DNA sequence.
- 5. The vector of Claim 2 wherein said vector further includes at least a portion of the adenoviral E3 DNA sequence.
- 6. A producer cell for generating adenoviral vector particles, said producer cell including an adenoviral E1 DNA sequence and an adenoviral E4 DNA sequence.
- 7. A method of generating infectious adenoviral particles, comprising:

transfecting the producer cell of Claim 6 with the adenoviral vector of Claim 3.

- 8. Infectious adenoviral particles generated by the method of Claim 7.
- 9. A method of effecting a gene therapy treatment, comprising:

administering to a host the infectious adenoviral particles of Claim 8 in an amount effective to produce a therapeutic effect in a host.

10. Bukaryotic cells transfected with the adenoviral particles of Claim 8.

WO 96/18418 PCT/US95/15947

11. The adenoviral vector of Claim 1 wherein said vector includes an adenoviral 5' ITR; an adenoviral 3' ITR; an adenoviral encapsidation signal; at least one DNA sequence encoding a protein or polypeptide of interest; and a promoter controlling the DNA sequence(s) encoding a protein(s) or polypeptide(s) of interest, said adenoviral vector being free of all or a portion of the adenoviral B1 and free of all or a portion of B2 DNA sequences.

- 12. The vector of Claim 11 wherein said vector is free of the adenoviral Ela and Elb DNA sequence.
- 13. The vector of Claim 12 wherein said vector is free of all or a portion of the adenoviral E2a DNA sequence.
- 14. The vector of Claim 11 wherein said vector is free of all or a portion of the adenoviral E2b DNA sequence.
- 15. The vector of Claim 11 wherein said vector further includes all or a portion of the adenoviral E3 DNA sequence.
- 16. A producer cell for generating adenoviral vector particles, said producer cell including an adenoviral E1 DNA sequence and an adenoviral E2a DNA sequence.
- 17. A method of generating infectious adenoviral particles comprising:

transfecting the producer cell of Claim 16 with the adenoviral vector of Claim 13.

- 18. Infectious adenoviral particles generated by the method of Claim 17.
- 19. A method of effecting a gene therapy treatment comprising:

administering to a host the infectious adenoviral particles of Claim 18 in an amount effective to produce a therapeutic effect in a host.

- 20. Bukaryotic cells transfected with the adenoviral particles of Claim 18.
- 21. The adenoviral vector of Claim 1 wherein said vector includes an adenoviral 5' ITR; and adenoviral 3' ITR; an adenoviral encapsidation signal; at least one DNA sequence

WO 96/18418 PCT/US95/15947

encoding a protein or polypeptide of interest; and a promoter controlling the DNA sequence(s) encoding a protein(s) or polypeptide(s) of interest, said adenoviral vector being free of all or a portion of each of the adenoviral E1, E2, and E4 DNA sequences.

- 22. The vector of Claim 21 wherein said vector is free of all or a portion of the adenoviral E2a DNA sequence.
- 23. The vector of Claim 21 wherein said vector is free of all or a portion of the adenoviral E2b DNA sequence.
- 24. The vector of Claim 21 wherein said vector further includes all or a portion of the adenoviral E3 DNA sequence.
- 25. A producer cell for generating adenoviral vector particles, said producer cell including an adenoviral E1 DNA sequence, an adenoviral E2a DNA sequence, and an adenoviral E4 DNA sequence.
- 26. A method of generating infectious adenoviral particles, comprising:

transfecting the producer cell of Claim 25 with the adenoviral vector of Claim 22.

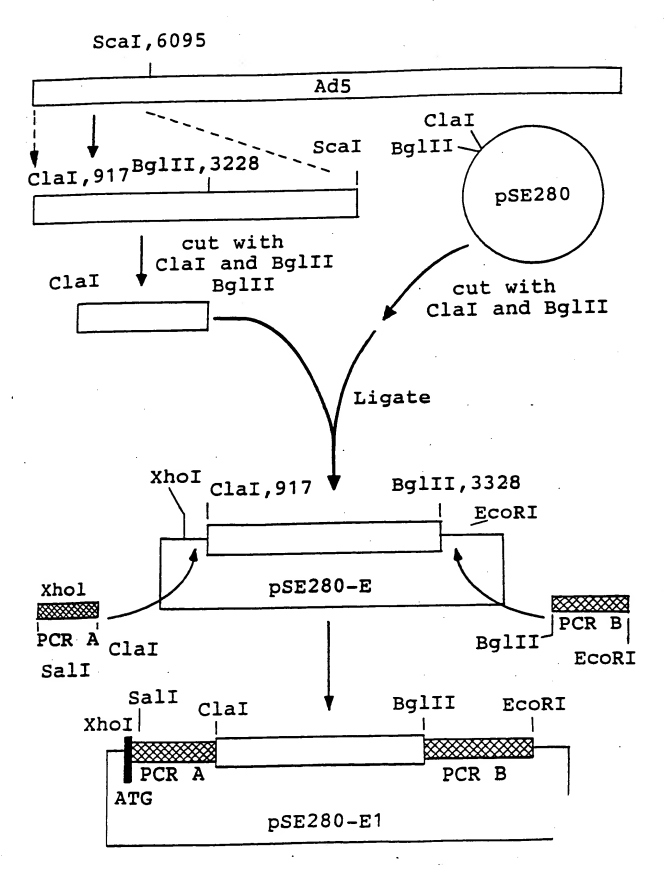
- 27. Infectious adenoviral particles generated by the method of Claim 26.
- 28. A method of effecting a gene therapy treatment, comprising:

administering to a host the infectious adenoviral particles of Claim 27 in an amount effective to produce a therapeutic effect in a host.

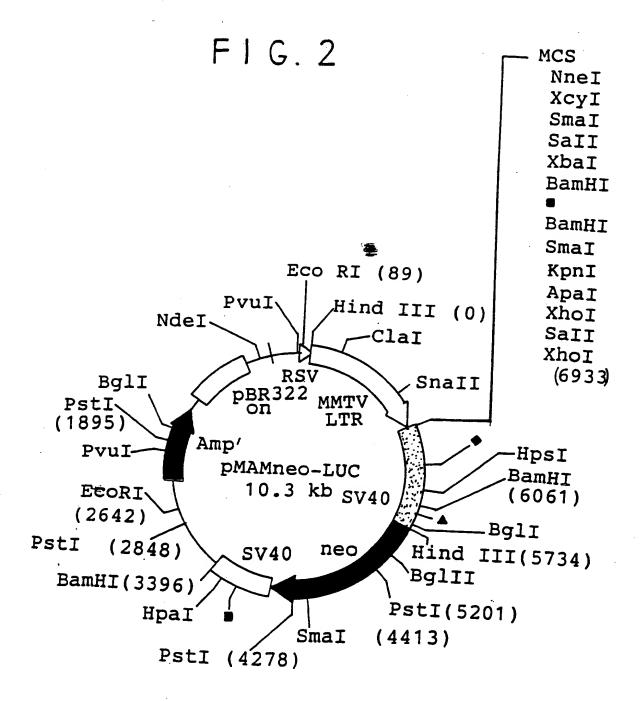
29. Rukaryotic cells transfected with the adenoviral vector of Claim 27.



FIG. 1

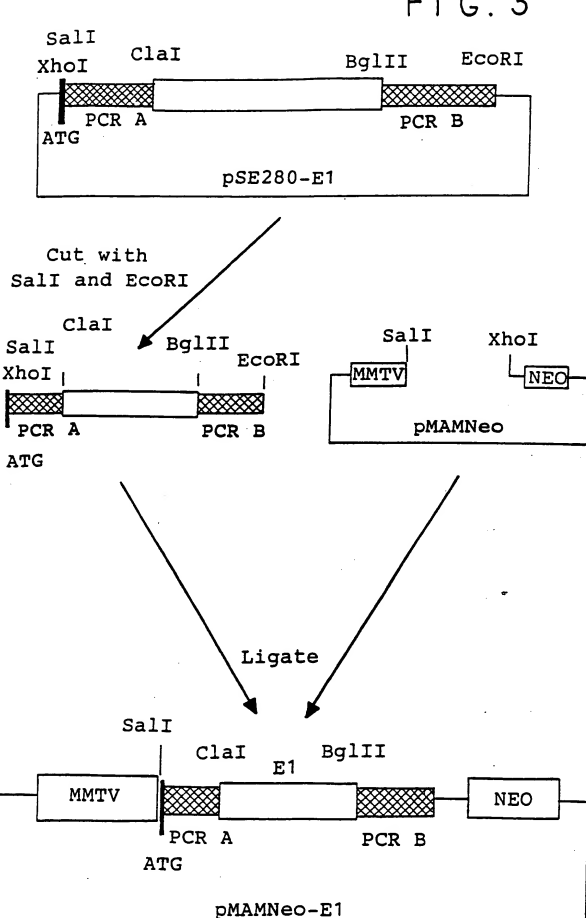


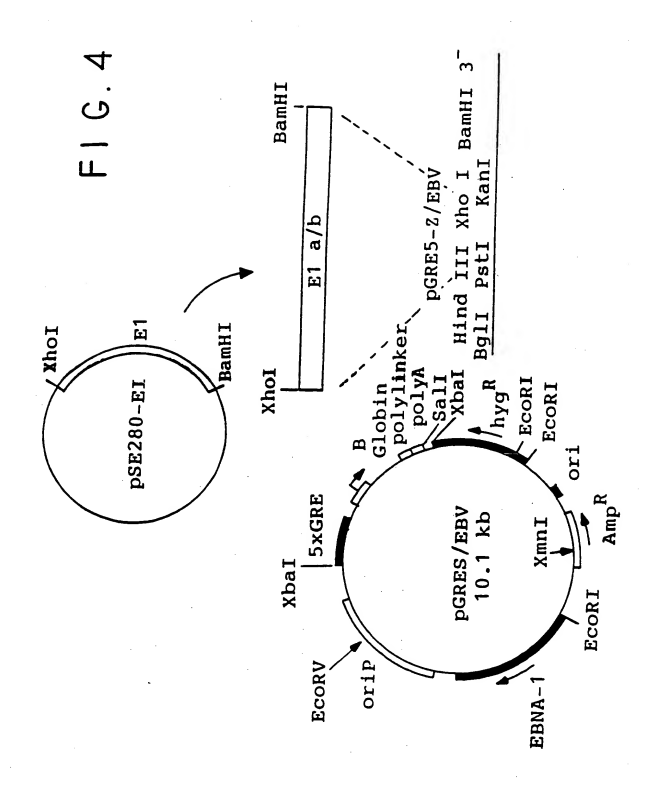
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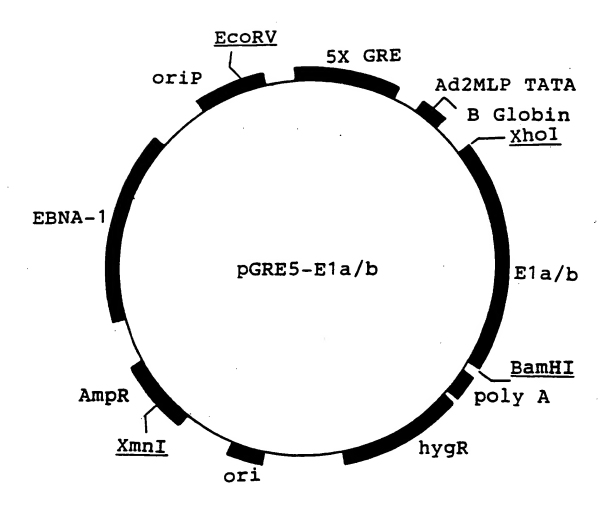
- luc pene
 - ▲ SV40 on & early promoter
 - SV40 early splicing region & polyadenylation





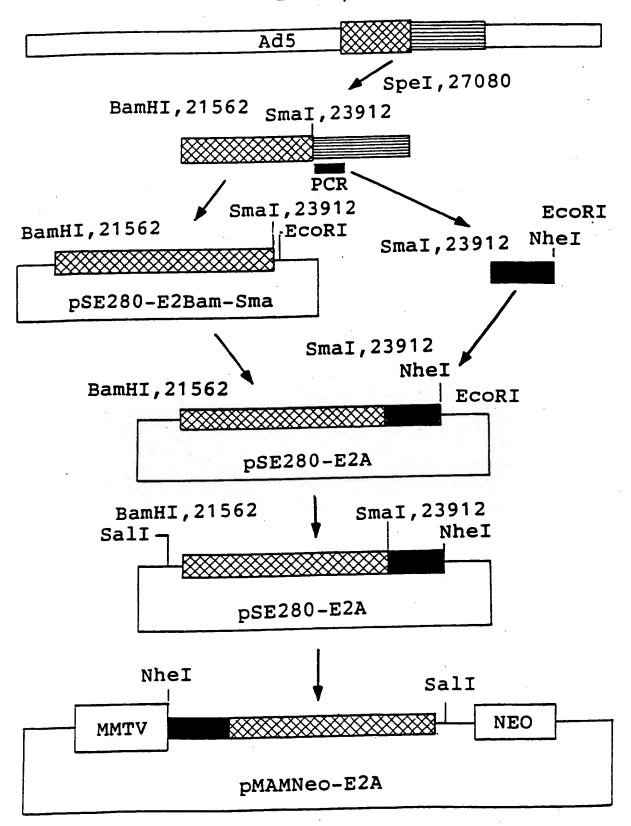






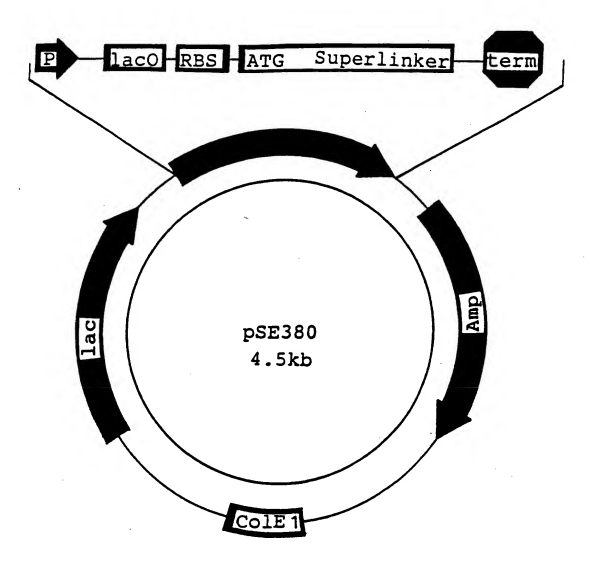
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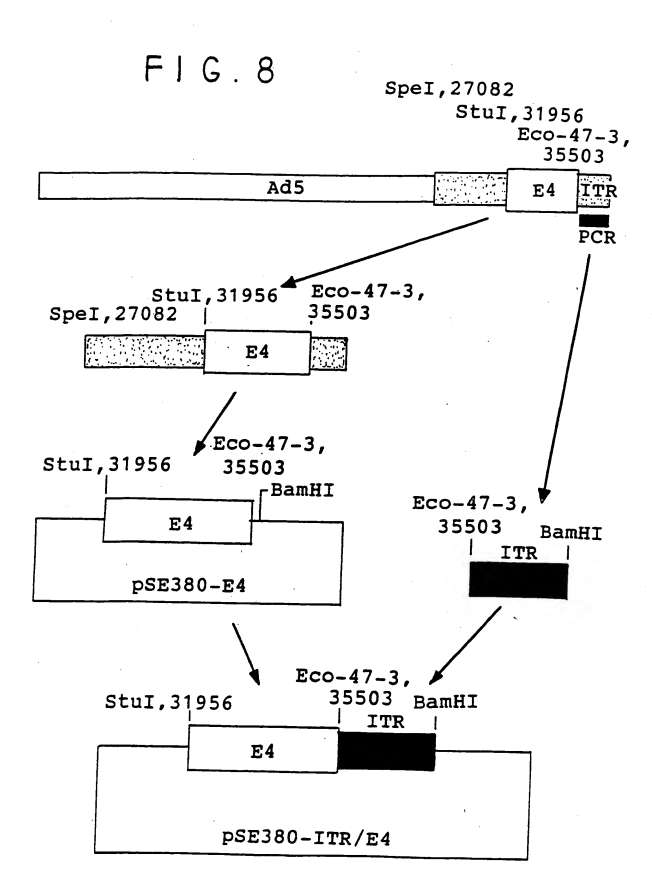
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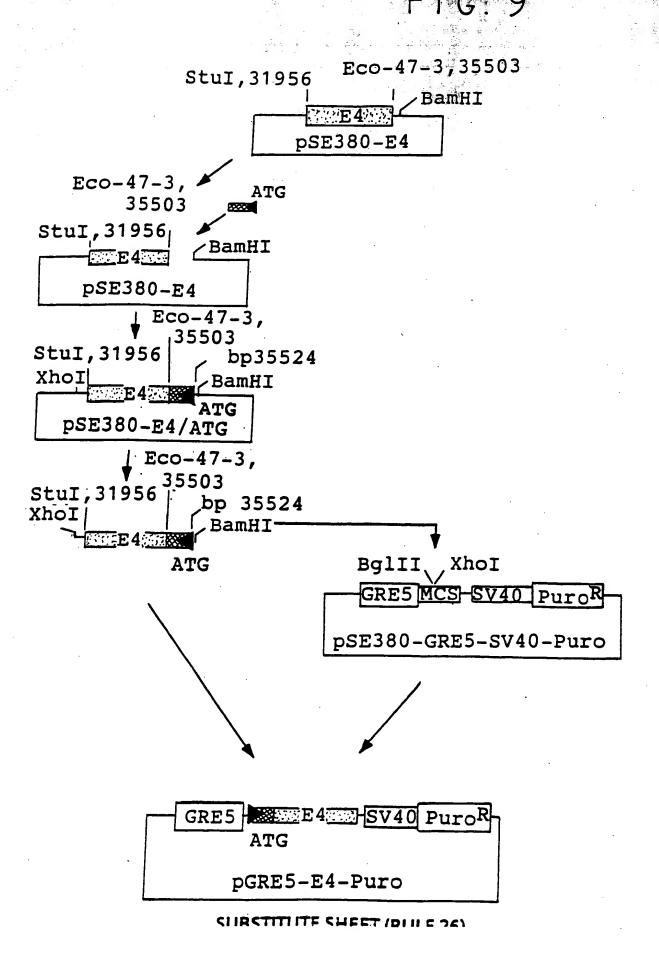
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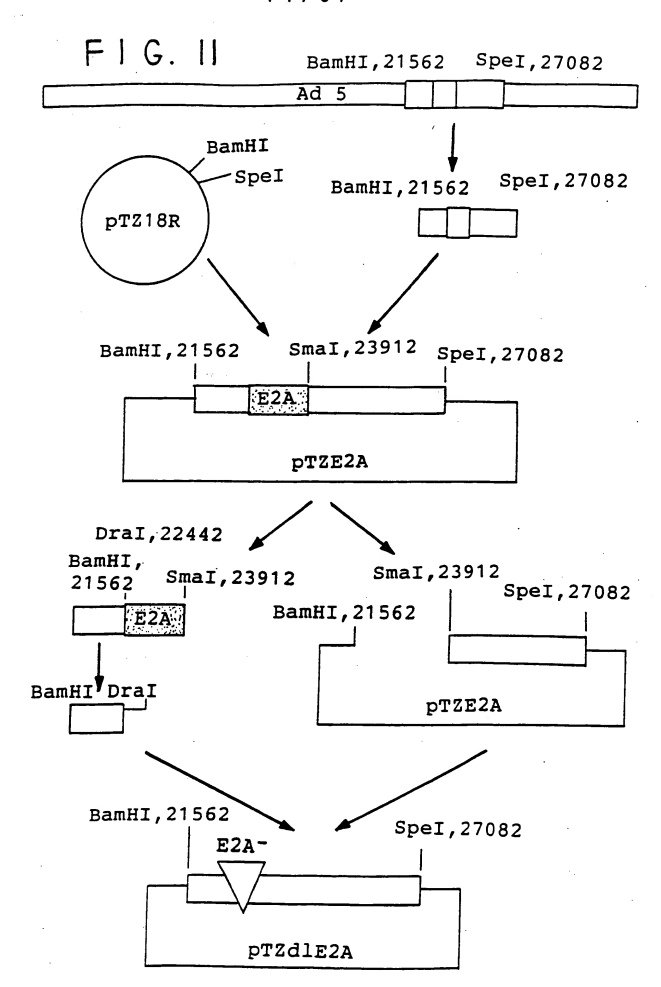




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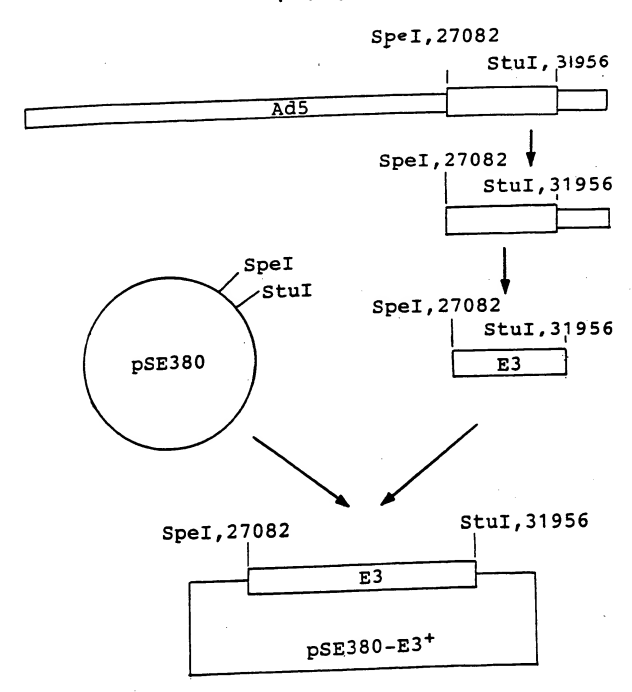




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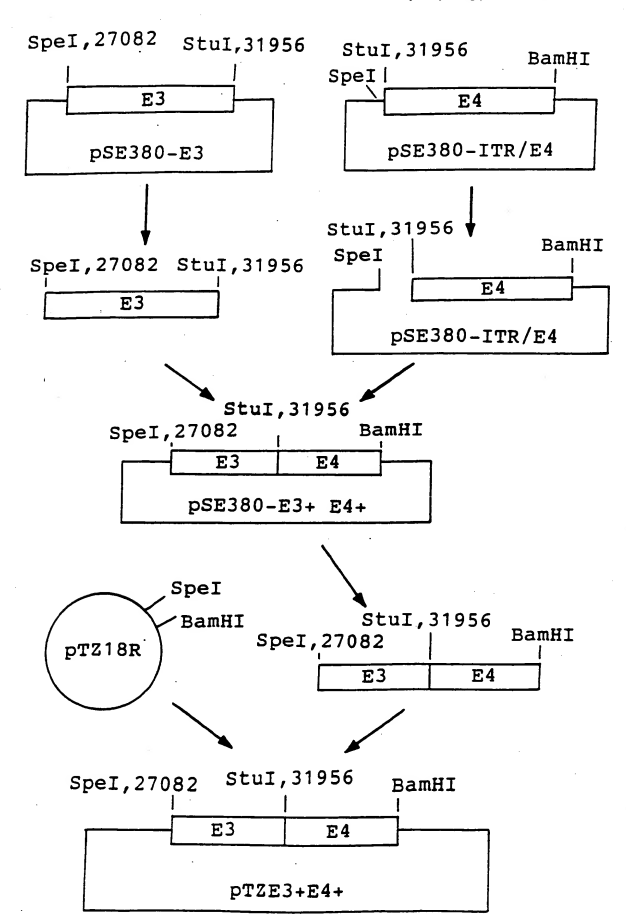


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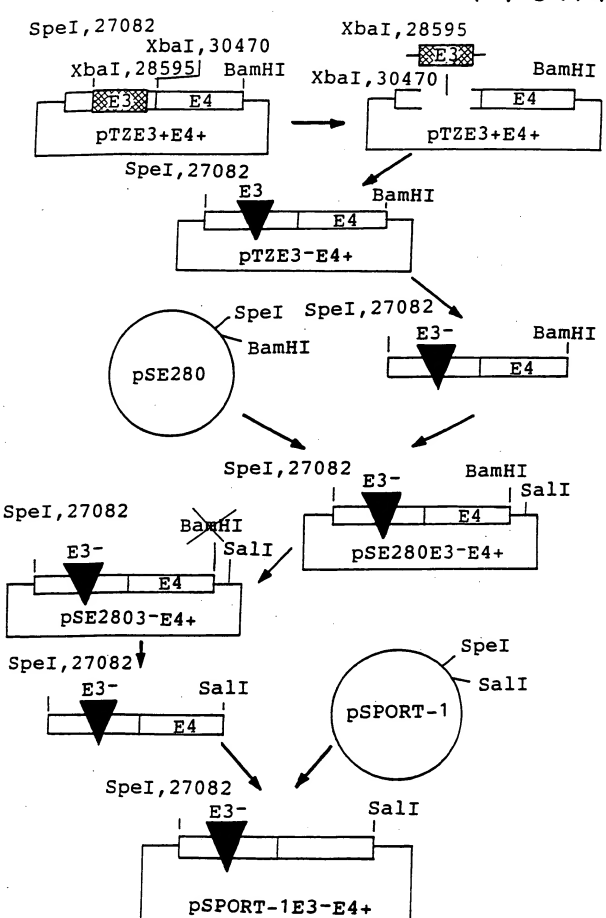


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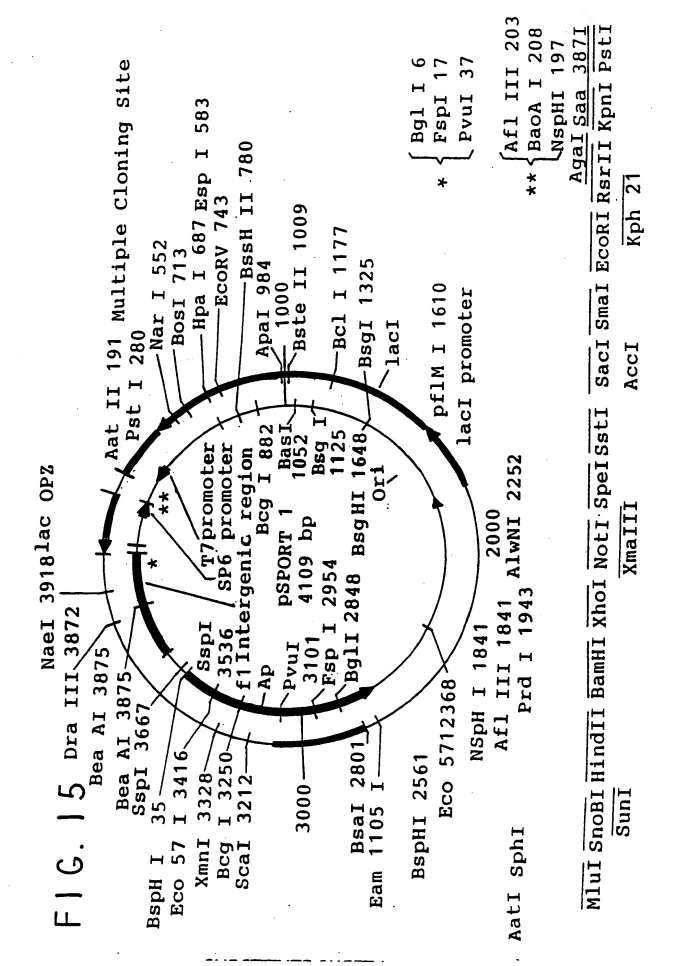


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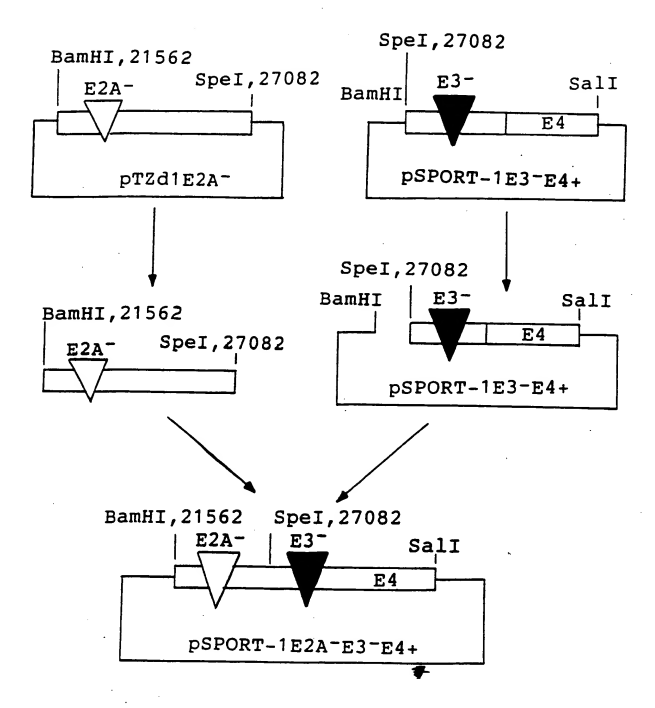
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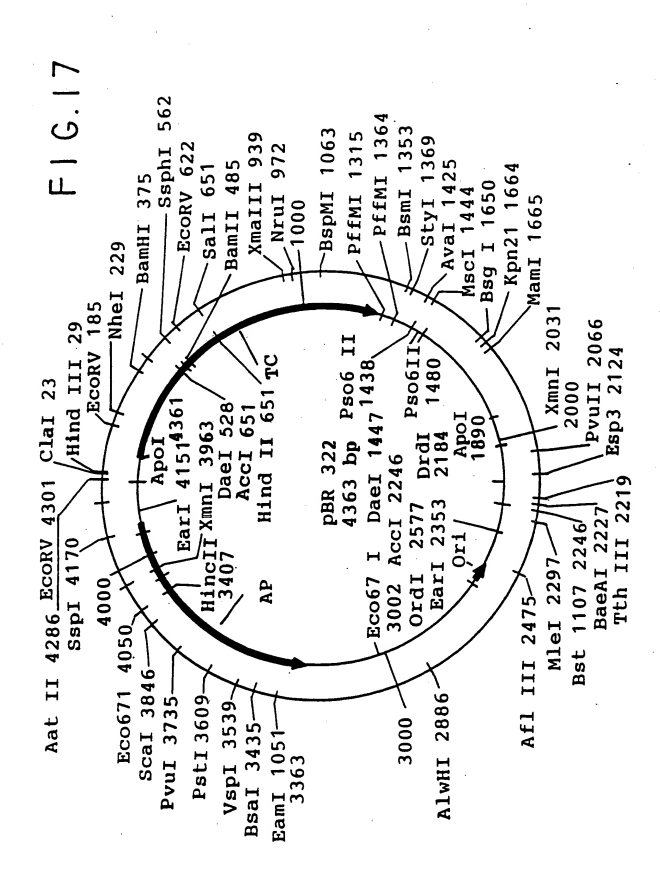


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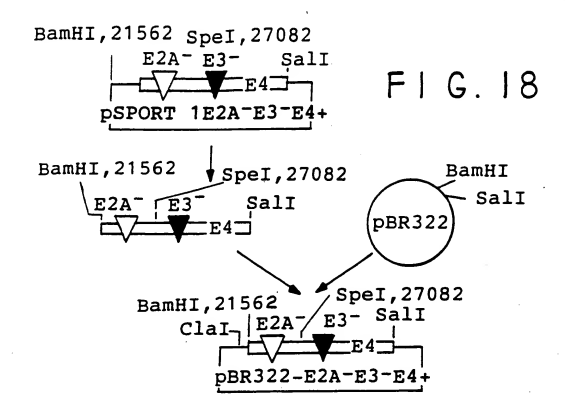


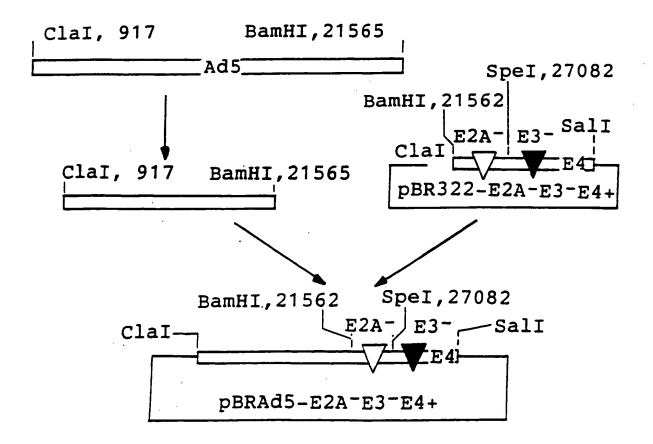


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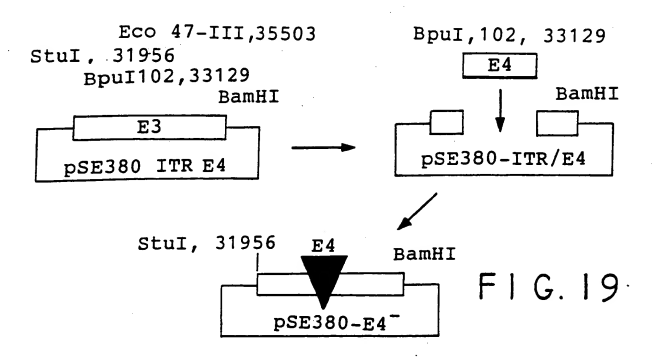
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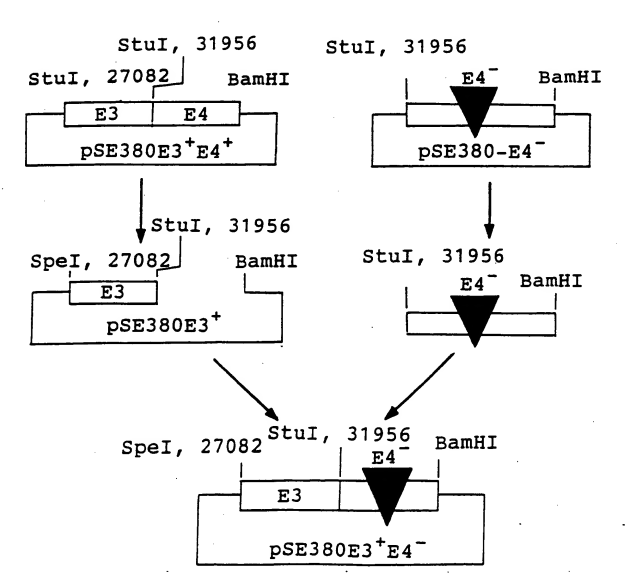












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FIG. 20

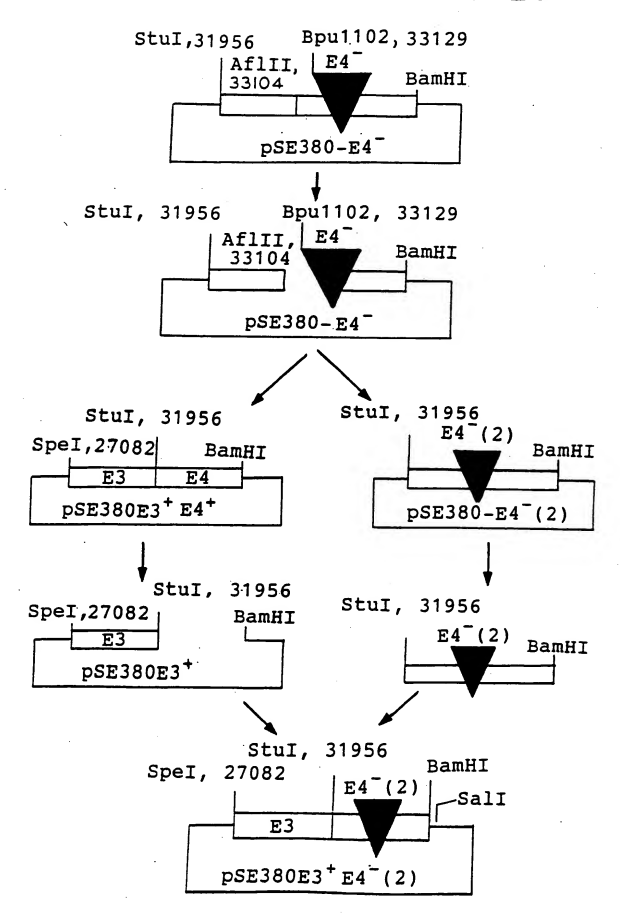
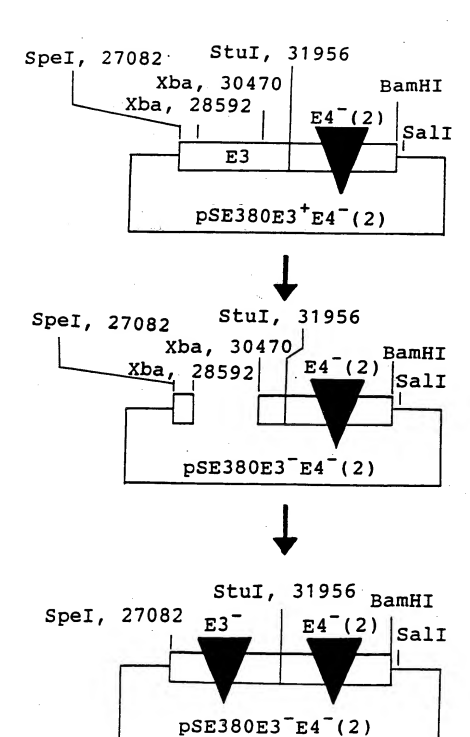




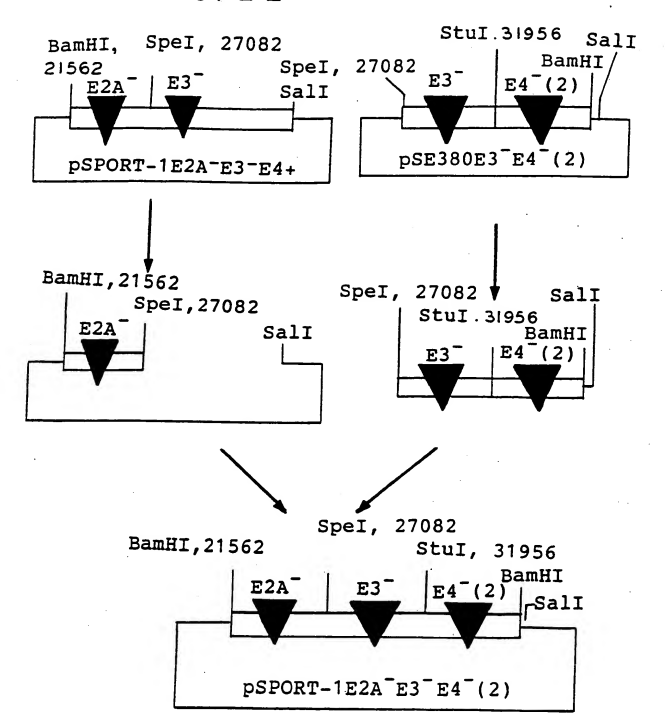
FIG. 21



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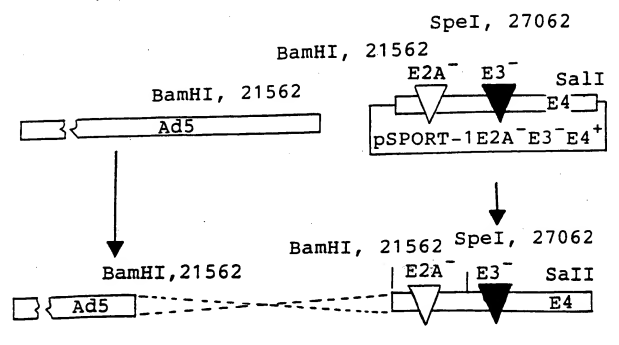
FIG. 22



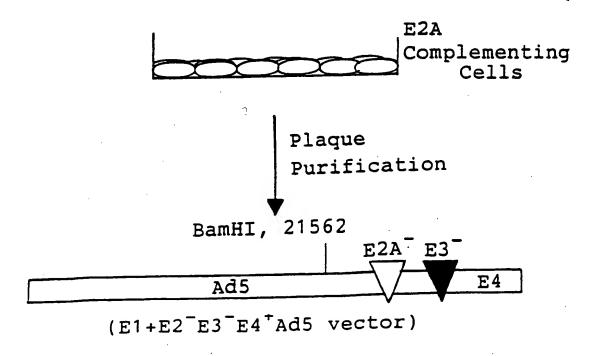
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FIG. 23

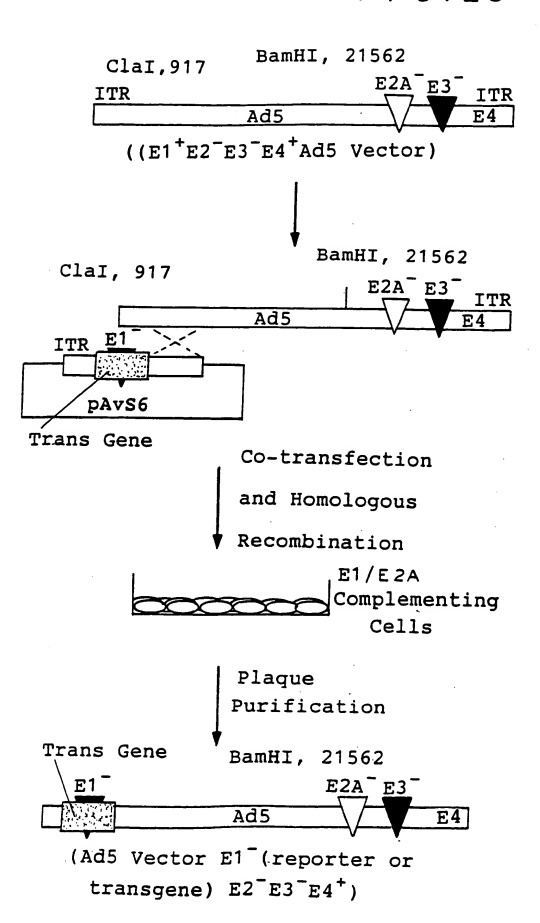


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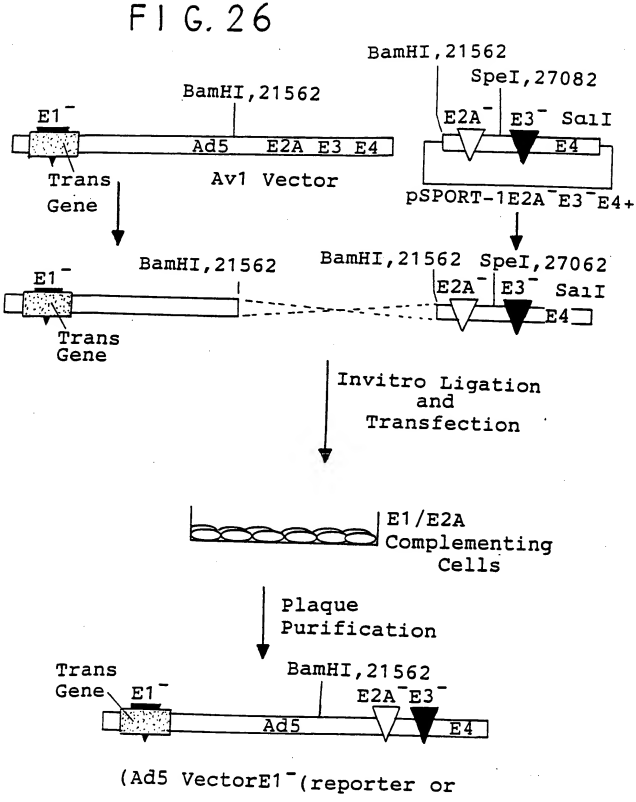


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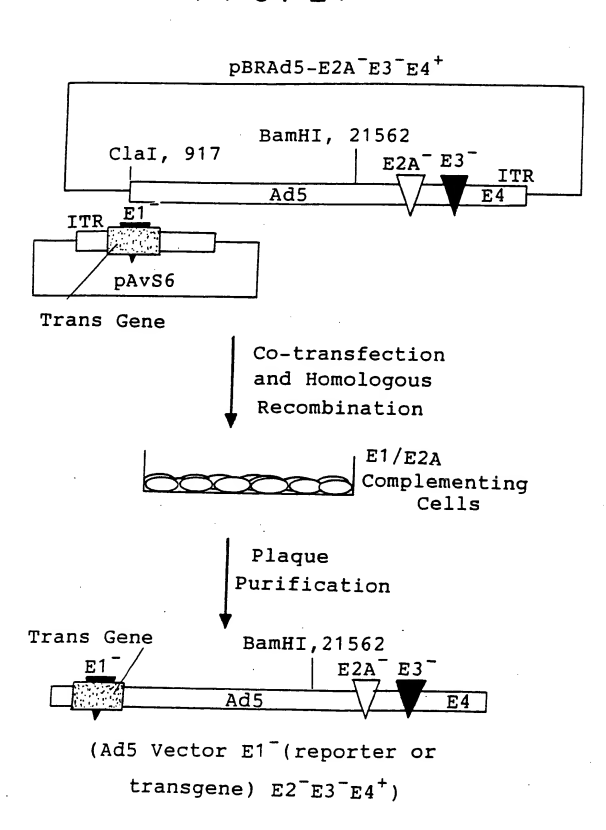




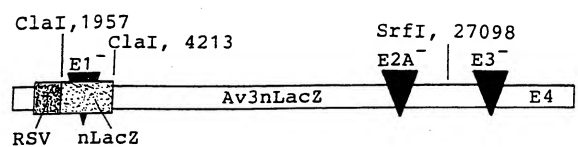
transgene) E2 E3 E4+)



FIG. 27







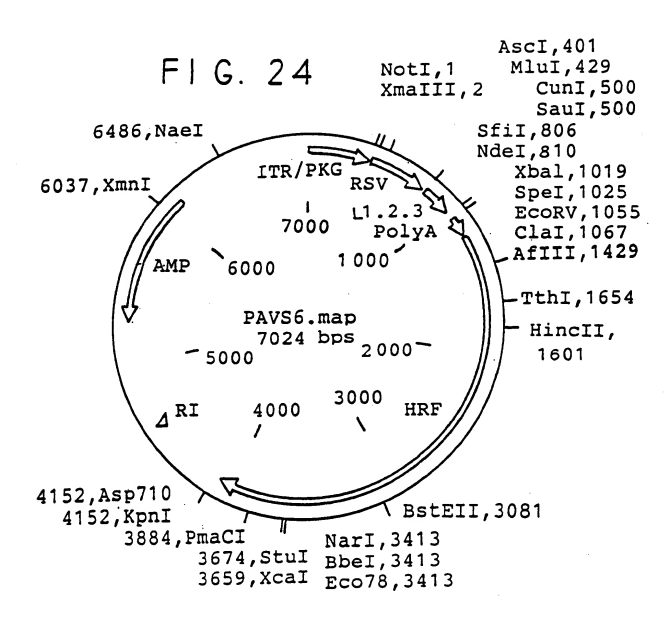
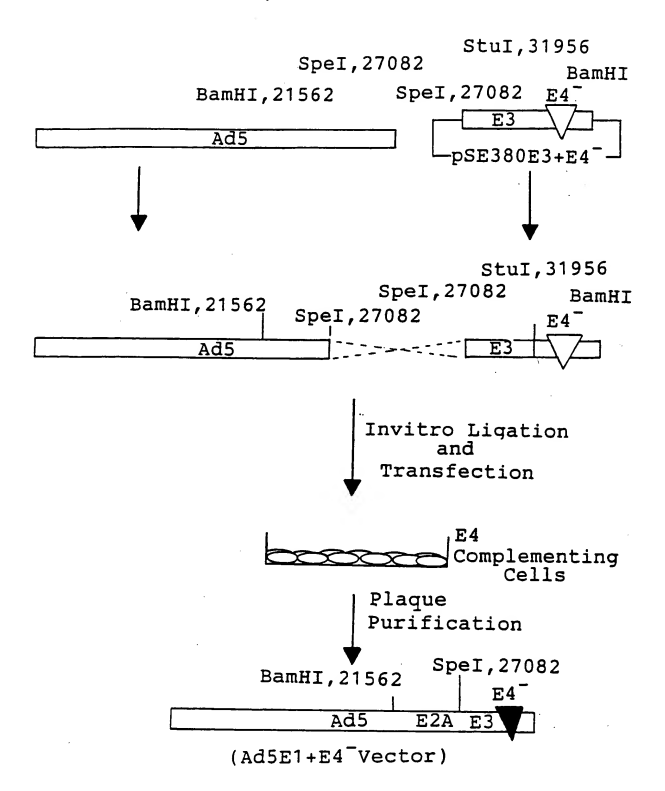
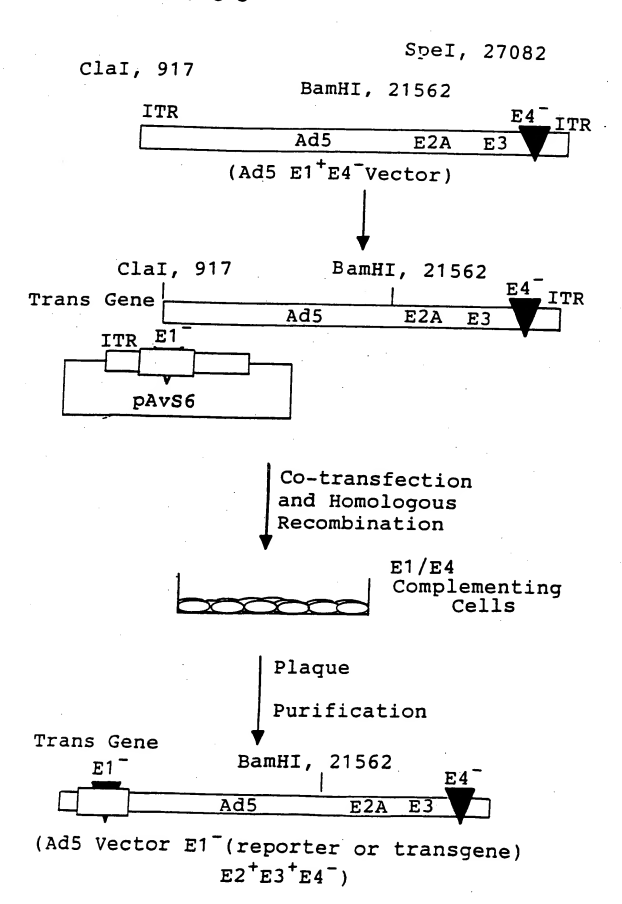




FIG. 29



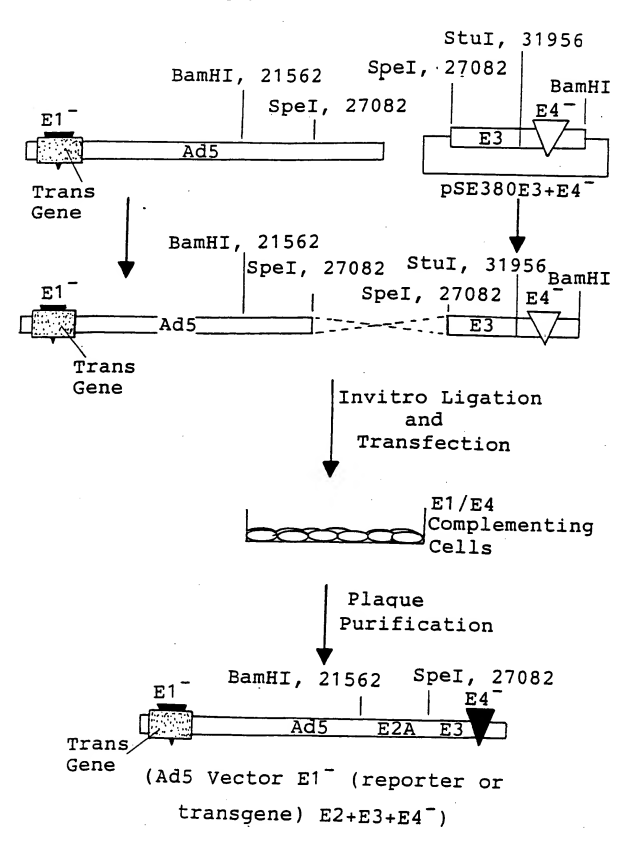
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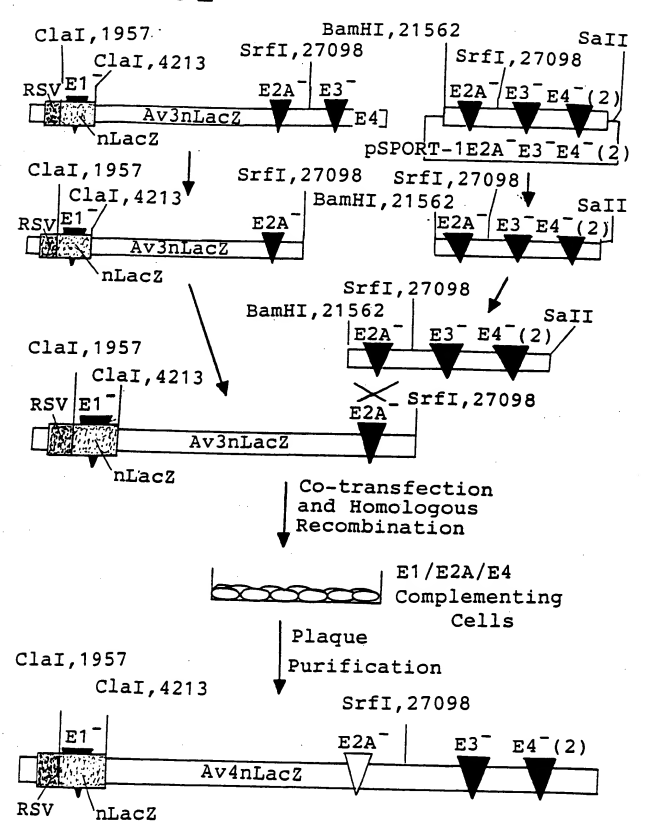
FIG. 31



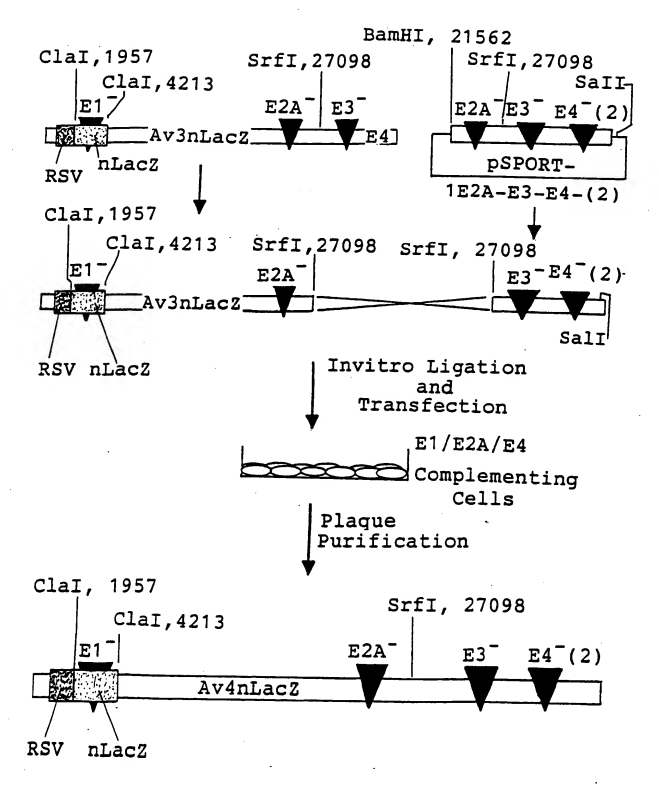
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F1G. 32



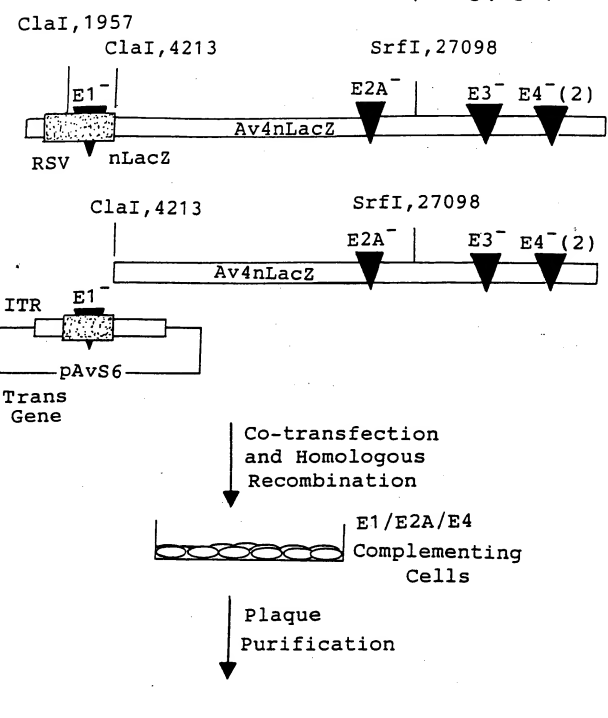


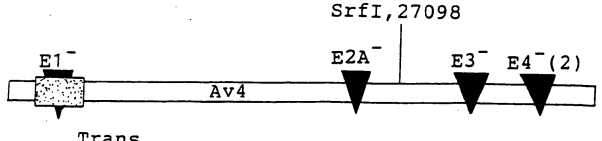


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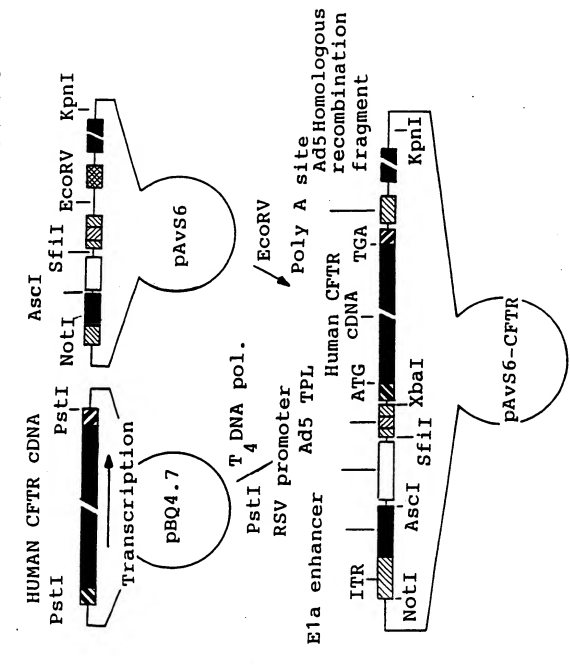






Trans Gene

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Adenovirus Vector Hexon Expression in A549 Cells

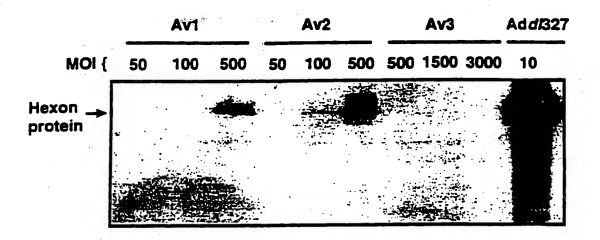
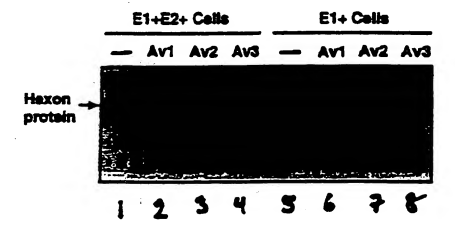


Fig. 36





Adenovirus Vector Hexon Expression



Figure



37/37

Adenovirus Vector DNA Replication

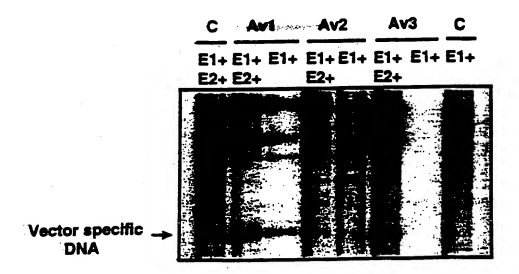


Fig.38

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A. CLASSIFICATION OF SUBJECT MATTER IPC(6) :A61K 48/00; C12N 15/00 US CL :514/44; 435/320.1 According to International Patent Classification (IPC) or to both national classification and IPC	
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B. FIELDS SEARCHED Minimum documentation searched (classification system followed by classification symbols)	
U.S. : 514/44; 435/320.1	
Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched	
Electronic data base consulted during the international search (name of data base and, where practicable, search terms used) APS, MEDLINE, EMBASE, BIOSIS, CAPLUS	
C. DOCUMENTS CONSIDERED TO BE RELEVANT	
Category* Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No.	
X WO, A, 94/28152 (IMLER ET AL) 08 DECEMBER 1994, see 1 2-29	
WO, A, 94/12649 (GREGORY ET AL) 09 JUNE 1994, see 1-29 entire document.	
PROCEEDINGS OF THE NATIONAL ACADEMY OF SCIENCES USA, Volume 91, issued June 1994, Engelhardt et al, "Ablation of E2A in recombinant adenoviruses improves transgene persistence and decreases inflammatory response in mouse liver", pages 6196-6200, see entire document.	
Purther documents are listed in the continuation of Box C. See patent family annex.]
Special categories of cited documents: "I" Inter-document published after the interactional filling date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention.	1
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the priority date chained Date of the actual completion f the international search Date of mailing f the international search report	+
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444

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